FINAL

Corrective Action Plan for the Risk-Based Remediation of the Military Gas Station Eglin Air Force Base Florida



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and

Eglin Air Force Base, Florida

March 1999

20000829 071

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FINAL

CORRECTIVE ACTION PLAN FOR THE RISK-BASED REMEDIATION OF THE MILITARY GAS STATION EGLIN AIR FORCE BASE, FLORIDA

AETC Contract No. F41689-96-D-0710 Order No. 5015

Prepared for
AIR FORCE CENTER FOR ENVIRONMENTAL EXCELLENCE
TECHNOLOGY TRANSFER DIVISION
BROOKS AIR FORCE BASE, TEXAS

and

EGLIN AIR FORCE BASE, FLORIDA

March 1999

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ACRONYMS AND ABBREVIATIONS

°C degrees centigrade
°F degrees Fahrenheit

AAS air sparge

AETC Air Education and Training Command

AFB Air Force Base

AFCEE/ERT Air Force Center for Environmental Excellence, Technology

Transfer Division

bgs below ground surface

BTEX benzene, toluene, ethylbenzene, and xylenes

CAP Corrective Action Plan

CAR Contamination Assessment Report

CH₄ methane

CO₂ carbon dioxide

COPC chemicals of potential concern

CSM conceptual site model DO dissolved oxygen EDB ethylene dibromide

FAC Florida Administrative Code

Fe2+ ferrous iron Fe3+ ferric iron

FDEP Florida Department of Environmental Protection

ft/day feet per day ft/ft feet per foot ft/yr feet per year

LCS laboratory control samples LTM long-term monitoring

LNAPL light, non-aqueous phase liquid

MDL method detection limit mg/L milligrams per liter msl mean sea level

MS/MSD matrix spike / matrix spike duplicate

MTBE methyl tert-butyl ether

mV millivolt

MW monitoring well

N nitrogen

NFA No-Further-Action

O₂ oxygen

ORP oxidation/reduction potential

OSHA Occupational Safety and Health Administration

OVA organic vapor analyzer OVM organic vapor meter

PAH polynuclear aromatic hydrocarbon

Parsons ES Parsons Engineering Science, Inc.

PEL permissible exposure limits

POC point of compliance

ppmv parts per million, volume per volume

PQL practical quantitation limit

QA quality assurance QC quality control RAP remedial action plan

RNA remediation by natural attenuation

SAP sampling and analysis plan

SB soil boring

SQL sample quantitation limit SVE soil vapor extraction

Tbsp tablespoon

TCLs target cleanup levels TDS total dissolved solids

TWA-TLV time-weighted average threshold limit value

TOC total organic carbon

TPH total petroleum hydrocarbons

TRPH total recoverable petroleum hydrocarbons

USEPA United States Environmental Protection Agency

UST underground storage tank

SECTION 1

INTRODUCTION

Parsons Engineering Science, Inc. (Parsons ES) was retained by the Air Force Center for Environmental Excellence, Technology Transfer Division (AFCEE/ERT) under Air Education and Training Command (AETC) Contract No. F41689-96-D-0710, Order No. 5015 to prepare a corrective action plan (CAP) to support a risk-based remediation decision for contaminated soil and groundwater at the Military Gas Station at Eglin Air Force Base (AFB) in Florida.

1.1 DESCRIPTION OF THE RISK-BASED APPROACH

The objective of risk-based remediation is to reduce the risk of specific chemicals to human health and/or ecological receptors such as animals or plant life. For any chemical to pose a risk, four elements must exist at the site:

- A source of chemical contamination that exceeds or could generate chemical contamination above health-protective or aesthetic standards;
- A mechanism of contaminant release;
- A human or ecological receptor available for chemical contact; and
- A completed pathway through which that receptor will contact the chemical.

If any one of these four elements is absent at a site, there is no current risk. The reduction or elimination of risk can be accomplished by limiting or removing any one of these four elements from the site.

The goal of this risk-based remediation approach is to find the most cost-effective method of reducing present and future risk by combining three risk reduction techniques:

- Chemical Source Reduction Achieved by natural attenuation processes over time or by engineered removals such as free product recovery, soil vapor extraction (SVE), or *in situ* bioventing.
- Chemical Migration Control Examples include natural attenuation of a groundwater plume, and SVE to prevent migration of hazardous vapors to a receptor exposure point.

 Receptor Restriction - Land use controls and site fencing to eliminate chemical exposure until natural attenuation and/or engineered remediation reduce the chemical source and/or eliminate the potential for chemical migration to an exposure point.

1.2 RISK-BASED APPROACH TASKS

The major tasks of this risk-based project are:

- Assessing available data and collecting any supplemental site characterization data necessary to define the nature, magnitude, and extent of soil and groundwater contamination and to document to what degree natural attenuation processes are operating at the selected sites;
- Determining whether an unacceptable risk to human health or the environment currently exists or may exist in the foreseeable future using applicable Florida Department of Environmental Protection (FDEP) guidance and regulations, contaminant fate and transport predictions, and exposure concentration estimates;
- Evaluating and recommending a remedial alternative that both reduces the source of contamination and minimizes or eliminates risks to potential receptors; and
- Documenting the remedial action selection process in a report that satisfies FDEP requirements.

1.3 REGULATORY REQUIREMENTS

This section describes Florida's tiered approach for risk-based remedial action at sites contaminated with petroleum products. The *Petroleum Contamination Site Cleanup Criteria* rule [Chapter 62.770 of the Florida Administrative Code (FAC)] (FDEP, 1997) presents guidance for determination of remedial requirements for closure of petroleum-contaminated sites, including several mechanisms for determining matrix-specific cleanup criteria. The regulations allow closure of petroleum release sites under several different scenarios, including:

- No-Further-Action (NFA) Proposal Without Conditions,
- · NFA Proposal With Conditions, or
- Monitoring-Only Proposal for Natural Attenuation.

A Remedial Action Plan (RAP) must be prepared for sites that do not meet the requirements for NFA or Natural Attenuation. Closure of a site under the NFA-Without-Conditions alternative would allow unrestricted future use of the site (e.g., residential land use), and therefore the requirements and allowable contaminant levels under this alternative are the most restrictive. The NFA-With-Conditions alternative requires that appropriate institutional or engineering controls be implemented to limit receptor exposure; sites seeking closure under this alternative are subject to potentially less stringent cleanup levels. A Natural Attenuation Monitoring Program is a

recognized means of remediating a site, with the goal of achieving the NFA cleanup target levels.

The actual or potential beneficial use of the groundwater and susceptibility of the aquifer to contamination are considered in the risk-based corrective action program to determine site-specific remediation target levels. All groundwater of the State of Florida is classified according to the following uses:

Class F-I: Potable water use: groundwater in a single source aquifer

described in Rule 62-520.460, FAC that has a total dissolved solids (TDS) content of less than 3,000 milligrams per liter (mg/L) and

was specifically reclassified as Class F-I by the Commission.

Class G-I: Potable water use: groundwater in a single-source aquifer that has

a TDS content of less than 3,000 mg/L.

Class G-II: Potable water use: groundwater in an aquifer that has a TDS

content of less than 10,000 mg/L, unless otherwise classified by

the Commission.

Class G-III: Non-potable water use: groundwater in an unconfined aquifer that

has a TDS content of 10,000 mg/L or greater; or that has a TDS content of 3,000-10,000 mg/L and either has been reclassified by the Commission as having no reasonable potential as a future source of drinking water, or has been designated by the FDEP as

an exempted aquifer pursuant to Rule 62-28.130(3), FAC.

Class G-IV: Non-potable water use: groundwater in a confined aquifer that has

a TDS content of 10,000 mg/L or greater.

The classification of the groundwater beneath the Military Gas Station is G-II (EA Environmental, Science, and Technology [EA], 1994).

1.3.1 No Further Action

Closure of a petroleum release site under a NFA Proposal (without or with conditions) requires that a site meet the following criteria:

- No free product is present (as specified in 62-770.680 (1)(a), FAC);
- No fire or explosion hazard is present due to release of petroleum or petroleum products;
- No "excessively contaminated soil" (as defined in 62-770.200, FAC) is present;
 and
- Matrix-specific target cleanup levels are met.

The Petroleum Contamination Site Cleanup Criteria rule (FDEP, 1997) incorporates matrix-specific Target Cleanup Levels (TCLs) for petroleum constituents in the form of

"look-up" tables or through reference to other applicable regulations (i.e., state groundwater or surface water regulations). Contaminant concentrations in all affected media at a site must be below all applicable TCLs for the site to qualify for a NFA (with or without conditions) proposal. However, the rule also allows for the development of alternative cleanup standards based on a site-specific risk assessment for use in a NFA Proposal with conditions. These site-specific alternative cleanup standards can be used in place of those presented in the look-up tables.

1.3.2 Natural Attenuation With Monitoring

The FDEP recognizes natural attenuation with monitoring as a viable site remediation strategy. The following criteria must be met to demonstrate that this strategy is appropriate for a site:

- No free product is present (as specified in 62-770.690 (1)(a),FAC);
- Contaminated soil is not present to the extent that it may increase cleanup costs;
- Groundwater contaminant concentrations above applicable target cleanup levels are not migrating beyond a temporary point of compliance (POC);
- Available data show an overall decrease in the mass of contamination; and
- Contaminant concentrations in groundwater do not exceed appropriate criteria (Table IX levels, 62-770, FAC); or the technical evaluations (as specified in 62-770.690 (1)(f), FAC) indicate that natural attenuation is an appropriate remedial alternative.

Natural attenuation with monitoring requires the establishment of a temporary POC based on site-specific conditions relating to land and groundwater use, potentially exposed populations, hydrogeology, and type and concentrations of contaminants. Concentrations of petroleum compounds at the POC cannot exceed levels presented in the rule. Monitoring of natural attenuation to show plume stability and/or contaminant reductions can eventually lead to a proposal for NFA With or Without Conditions.

1.3.3 Summary

In summary, the *Petroleum Contamination Site Cleanup Criteria* rule allows a rapid determination of whether or not a site can qualify for a NFA proposal and/or the appropriateness of natural attenuation with monitoring as a remedial strategy. The rule allows for inclusion of site-specific information in developing alternative cleanup levels for NFA with conditions, and provides guidance on preparation of a RAP if active remediation is warranted.

1.4 REPORT ORGANIZATION

This CAP consists of eight sections, including this introduction, and five appendices. Site background, including operating history and a review of environmental site investigations conducted to date, is provided in the remainder of this section. Section 2 summarizes the 1998 site characterization activities performed by Parsons ES. Physical

characteristics of the Military Gas Station and surrounding area are described in Section 3. A Tier 1 evaluation is completed in Section 4 to identify those site contaminants that are considered chemicals of potential concern (COPCs). Section 5 summarizes the nature and extent of COPC contamination at the site. Section 6 addresses the effects of natural chemical attenuation processes that are documented to be occurring at the site, and presents chemical fate and transport and receptor exposure analyses. The Tier 2 evaluation is detailed in Section 7. Section 8 presents a long-term monitoring plan (LTM). Section 9 presents references used in preparing this CAP.

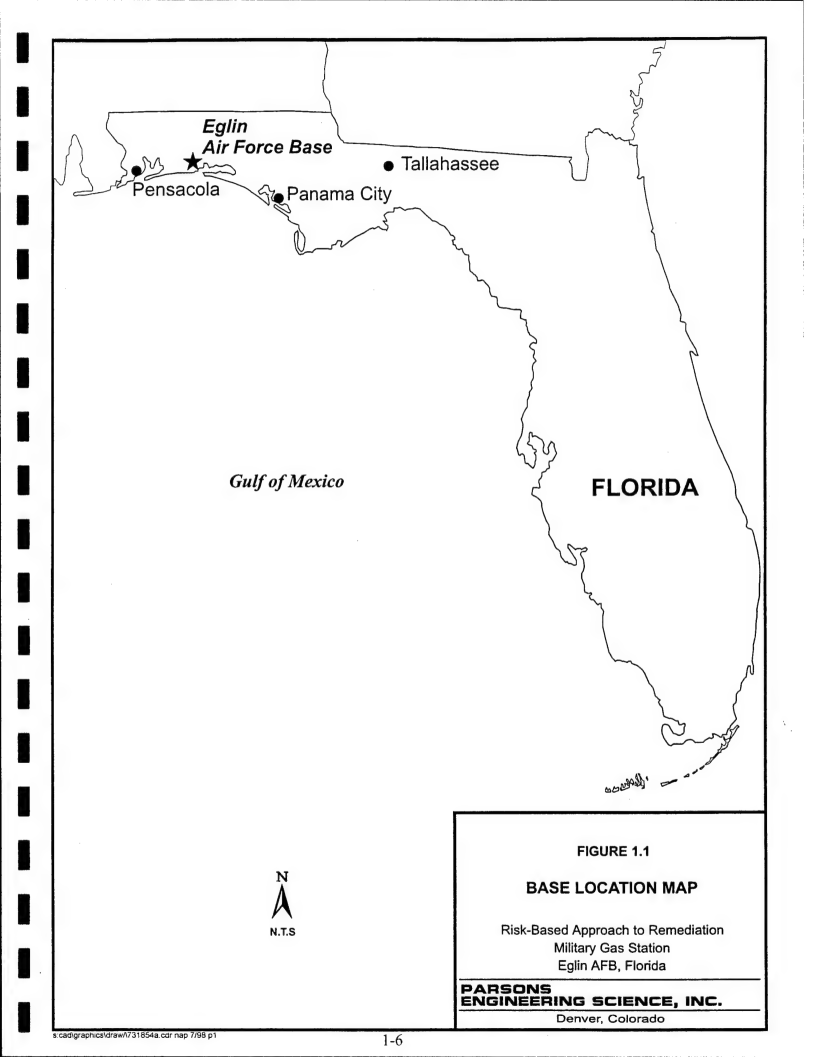
Analytical data sheets and chain-of-custody records are in Appendix A. Pertinent information from prior investigations is presented in Appendix B. Boring logs, groundwater sampling forms, and well construction diagrams for all sampling activities completed by Parsons ES during the March 1998 field effort are included in Appendix C. Appendix D includes the input and output from the aquifer slug test analyses. Appendix E includes the supporting documentation for the quantitative calculations used in the predictive chemical fate assessment.

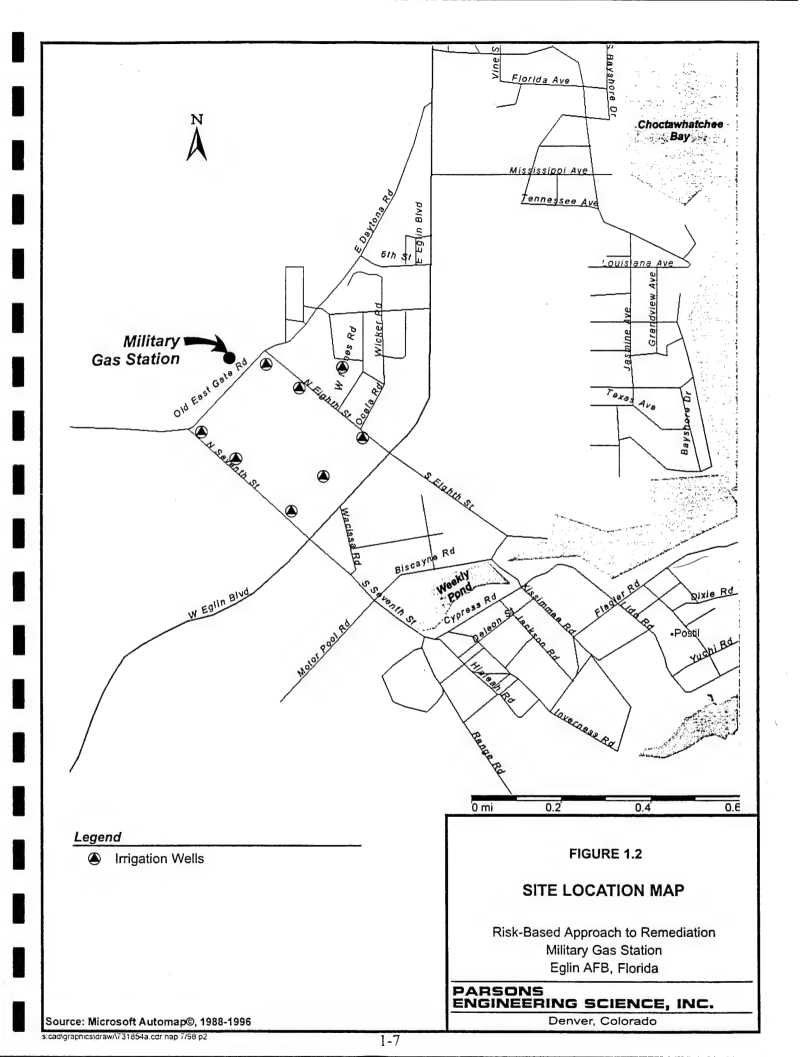
1.5 SITE DESCRIPTION AND BACKGROUND

Eglin AFB is located in the Florida panhandle on the coast of the Gulf of Mexico (Figure 1.1). The Military Gas Station is located at the intersection of Daytona Road and Okaloosa Avenue on the Eglin Main Base (Figure 1.2). The age of the gas station is unknown. Features of the site include a canopy sheltering 2 pump islands, an unmanned kiosk containing a computer system, and a tank field containing three 15,000-gallon fiberglass underground storage tanks (USTs) (Figure 1.3). Two of the tanks contain unleaded gasoline; the other tank contains diesel fuel.

Five USTs, which stored gasoline and diesel fuel, were previously located within the former tankfield (Figure 1.3). The USTs were removed in September 1991, and approximately 400 cubic yards of soil were excavated (EA, 1994b). During the tank closure assessment, the surrounding soil was screened for volatile hydrocarbons with an organic vapor analyzer (OVA). Soil in the vicinity of the south end of the tanks and near the fuel dispensers was classified as excessively contaminated [>50 parts per million, volume per volume (ppmv) for Mixed Product Analytical Group]. The contaminated soil was allowed to aerate before being returned to the excavation as fill.

The present tankfield and product piping were installed during September 1991. A Contamination Assessment Report (CAR) (EA, 1993a), CAR Addendum (EA, 1993b), and Supplemental CAR (EA, 1994a) have been completed for the site, and a RAP (EA, 1994b) has been approved. In support of the assessment and remedial action, 13 shallow monitoring wells (MWs), 2 deep MWs, 2 SVE wells, 1 air sparging (AAS) well, and 15 soil borings (SBs) were completed at the site, and an air sparging and SVE system was installed and has been operating for 2 to 3 years to remediate site soil and shallow groundwater in the source area (Parsons ES, 1997c).





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SECTION 2

SITE CHARACTERIZATION ACTIVITIES

Several soil and groundwater investigations have been conducted at the Military Gas Station. These investigations focused on characterizing and delineating dissolved hydrocarbons in groundwater and residual fuel hydrocarbons in soils. Parsons ES conducted an investigation at the site during March 1998 to collect site-specific data relevant to quantifying the effects of natural contaminant attenuation processes and to facilitate development and implementation of a risk-based remedial action for the Military Gas Station. Soil gas, soil, and groundwater were sampled to:

- Further delineate the extent of contamination;
- Assess temporal trends in soil and groundwater contaminant concentrations;
- · Support contaminant fate and transport analyses; and
- Develop appropriate exposure-point concentrations to compare to final remediation goals.

Data collected during previous investigations were used to augment this study. Emphasis was placed on collecting data documenting the natural biodegradation and attenuation of fuel hydrocarbons in soils and groundwater at the site.

The March 1998 supplemental site characterization activities performed by Parsons ES at the Military Gas Station are briefly described in the remainder of this section. Most site characterization procedures (i.e., soil, soil gas, and groundwater sampling procedures) are described in detail in the project Sampling and Analysis Plan (SAP) (Parsons ES, 1997a).

2.1 SCOPE OF DATA COLLECTION ACTIVITIES

As part of the risk-based remedial approach for the Military Gas Station, field data collection efforts focused on investigating specific chemical constituents that potentially pose a threat to human health or the environment. The chemicals targeted for study at this site were identified from previous site investigations and the chemical composition of the primary contaminant source (i.e., release(s) of gasoline from the former USTs). The petroleum hydrocarbons and associated constituents identified and addressed as part of this study, as either historically above FDEP levels or previously unquantified, include benzene, toluene, ethylbenzene, and xylenes (BTEX); methyl tertiary butyl ether (MTBE); ethylene dibromide (EDB); polynuclear aromatic hydrocarbons (PAHs);

total recoverable petroleum hydrocarbons (TRPH); and lead. These analytes were targeted based on previous site assessment results.

The risk-based investigation for the Military Gas Station was conducted according to the methodologies presented in the Work Plan for the Risk-Based Investigation and Closure of the Base Exchange Service Station and the Military Gas Station (Parsons ES, 1998), hereafter referred to as the work plan. The work plan was developed according to available guidelines and requirements of the FDEP to support site closure.

The following sampling and testing activities were performed by Parsons ES during March 1998 at the site as part of this investigation:

- Conducted slug tests at two existing monitoring wells;
- Drilled three soil borings;
- Collected five subsurface soil samples for fixed-base laboratory analysis from the three boreholes;
- Collected groundwater samples from four existing groundwater monitoring wells;
- Collected one soil gas sample for laboratory analysis.

Analytical method detection limit (MDL) requirements were considered before site characterization work was initiated. Suitable analytical methods and quality control (QC) procedures were selected (Parsons ES, 1997a) to ensure that the data collected under this program are of sufficient quality to be used in a quantitative risk assessment.

Soil and groundwater samples were analyzed in the field and by Quanterra, Inc. of Arvada, Colorado; Dallas, Texas; and Tampa, Florida. Soil gas samples were analyzed in the field and by Air Toxics, Ltd. of Folsom, California. The laboratory data sheets and chain-of-custody records are presented in Appendix A. The analytical protocol for all samples is summarized in Table 2.1. Table 2.2 summarizes the field and fixed-base laboratory analyses performed by sampling location. These analyses and measurements were performed for various inorganic, geochemical, and physical parameters to document natural biodegradation processes and to assess the potential effectiveness of low-cost source reduction technologies.

2.2 SUBSURFACE SOIL SAMPLING

Soil samples were collected adjacent to previously-drilled soil boreholes B, V, and K to obtain soil total organic carbon (TOC) data and to further characterize soil contamination at the site at locations where previous investigations indicated relatively high soil contaminant concentrations (Appendix B). The soil boring locations are presented on Figure 2.1. Soils were sampled to facilitate evaluation of the potential for contaminant partitioning from soil into groundwater and soil gas, and to assess the magnitude of any changes in contaminant concentrations that have occurred over time. These borings were advanced using a Geoprobe® hydraulic sampling rig as described in the SAP (Parsons ES, 1997a).

TABLE 2.1

ANALYTICAL PROTOCOL FOR

GROUNDWATER, SOIL, AND SOIL GAS SAMPLES

Military Gas Station

Eglin AFB, Florida

MATRIX	METHOD	WHERE ANALYZED
ROUNDWATER		
Ferrous Iron (Fe ⁺²)	Colorimetric, Hach Method 8146	Field
Sulfate (SO ₄ ⁻²)	Colorimetric, Hach Method 8051	Field
Conductivity	Direct reading meter	Field
Dissolved Oxygen	Direct reading meter	Field
pH	Direct reading meter	Field
Redox Potential	Direct reading meter	Field
Temperature	Direct reading meter	Field
BTEX	SW8020A	QUANTERRA*
EDB	SW8011 / Method 504	QUANTERRA
Polynuclear Aromatic Hydrocarbons	SW8310	QUANTERRA
Total Recoverable Petroleum Hydrocarbons (TRPH)	FL-PRO (C8-C40)	QUANTERRA
Methane (CH ₄)	RSK-175	QUANTERRA
Nitrate as Nitrogen (NO ₃ ⁻¹ -N)	E300.0	QUANTERRA
Lead	SW7421	QUANTERRA
OIL		
BTEX + MTBE	SW8020A	QUANTERRA
Polynuclear Aromatic Hydrocarbons	SW8310	QUANTERRA
Total Recoverable Petroleum Hydrocarbons (TRPH)	FL-PRO (C8-C40)	QUANTERRA
Total Organic Carbon	SW9060	QUANTERRA
OIL GAS		
BTEX	TO-3	Air Toxics ^{b/}
Total Petroleum Hydrocarbons (TPH)	TO-3	Air Toxics
Oxygen	Direct reading meter	Field
Carbon Dioxide	Direct reading meter	Field

Notes:

- a/ Quanterra, Inc. of Arvada, Colorado; Dallas, Texas (methane only); and Tampa, Florida (TRPH only).
- b/ Air Toxics LTD. of Folsom, California

TABLE 2.2 SAMPLE ANALYSES BY LOCATION

Military Gas Station Eglin AFB, Florida

			4	Egin AFB, Florida	Florida					
Sample Location	n B	Λ	Λ	¥	×	Mil SG1	EA-4	EA-5	EA-8	EA-10
Sample Matrix	Soil	Soil	Soil	Soil	Soil	Soil Gas	Water	Water	Water	Water
Sample Depth (ft. bgs)	() 25 - 27	8 - 10	26 - 28	8 - 10	15 - 17	NA	NA	NA	NA	٧N
ANALYTE										
BTEX	×		X		×		×	×	×	×
MTBE	×		×		×					
EDB							×	×	×	×
PAH	×		Х		×		×			×
TRPH	X		X		×		×			×
TOC		X		X						
Lead							×			×
Methane							×	×	X	×
ORP							×	×	×	×
Conductivity							×	×	×	×
Dissolved O ₂							×	×	×	×
Temperature							×	×	×	×
Hd							×	×	×	×
. Fe ²⁺							×	×	×	×
Sulfate	:						×	×	×	×
Nitrate							×	×	×	×
BTEX/TPH						X				
oxygen						X				
carbon dioxide						×				
Notes:										

Notes:

ft. bgs = Feet below ground surface

NA = Not applicable

BTEX = Benzene, Toluene, Ethylbenzene, and Xylenes

MTBE = Methyl Tert Butyl Ether

EDB = Ethylene dibromide

PAH = Polynuclear Aromatic Hydrocarbons

TRPH = Total Recoverable Petroleum Hydrocarbons

TOC = Total organic carbon

ORP = Oxidation-reduction potential

Fe²⁺ = Ferrous iron

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A total of five soil samples from the three boreholes were submitted to Quanterra, Inc. for laboratory analysis. Samples from all three boreholes were described for lithology and field screened for volatile organic vapors using a organic vapor meter (OVM). Soil borehole information is summarized in Table 2.3, and borehole logs are included in Appendix C. Soil analytical results are summarized and discussed in Sections 4 and 5.

2.3 GROUNDWATER SAMPLING

Groundwater samples were collected from four existing monitoring wells at the site in March 1998. The groundwater sampling locations are listed on Table 2.2 and shown on Figure 2.1. Samples collected from the four wells were analyzed for fuel-related contaminants and for various inorganic and geochemical indicators to evaluate natural chemical and physical attenuation processes that are occurring at the site. Field and laboratory analyses for each groundwater sampling location are summarized in Table 2.2.

All monitoring wells were purged using a positive displacement pump with dedicated polyvinyl chloride (PVC) tubing. Purging consisted of removing groundwater from the well until the pH, DO concentration, oxidation-reduction potential (ORP), conductivity, and temperature stabilized.

Within 24 hours of the purge event, groundwater samples were collected from the monitoring wells using dedicated teflon bailers. The water was carefully poured down the inner walls of each sample bottle to minimize aeration of the sample. Sample bottles for BTEX, MTBE, methane, and/or Hach® field analyses were filled so that there was no headspace or air bubbles within the container. One duplicate sample was collected during the groundwater sampling event.

Field and laboratory groundwater analytical results are discussed in Sections 4 and 5 of this report. These analytical results are used in Section 6 to evaluate the natural physical, chemical, and biological processes that are affecting the COPCs at this site.

2.4 SOIL GAS MEASUREMENTS

Soil gas sampling was performed at the site using both field (semi-quantitative) and fixed-base laboratory (quantitative) analyses. The purpose of soil gas sampling was to assess the potential risk to future workers at the site from inhalation of volatilized contaminants, and to determine whether or not sufficient oxygen (O₂) is available in the soil gas to sustain aerobic fuel hydrocarbon biodegradation. If O₂ concentrations are significantly lower than background values, and carbon dioxide (CO₂) concentrations are higher than background levels, then the occurrence of aerobic fuel hydrocarbon biodegradation can be inferred. In addition, the O₂ levels allow an assessment of whether there is sufficient O₂ to sustain continuing aerobic biodegradation without engineered addition of O₂ via *in situ* bioventing.

One soil gas sample was collected at the location Mil SG1 shown on Figure 2.1. The sample was screened using a field instrument to measure O₂ and CO₂, collected in a SUMMA® canister, and submitted to Air Toxics, Ltd. in Folsom, California for analysis of total petroleum hydrocarbons (TPH) and BTEX using US Environmental

TABLE 2.3 SOIL BORING SUMMARY

Military Gas Station Eglin AFB, Florida

Boring Location	Boring Date	Total Depth (ft bgs) ^{a/}
В	3/29/98	25.0
V	3/29-30/98	29.0
K	3/29/98	20.0

a/ ft bgs = feet below ground surface

Protection Agency (USEPA) Method TO-3. Analytical results for the soil gas sample are summarized in Sections 4 and 5.

2.5 SLUG TESTS AND ANALYSIS

Four slug tests were conducted at two existing monitoring wells at the Military Gas Station in March 1998. Two tests were conducted at each of wells EA-10 and EA-11. The data were analyzed using AQTESOLV aquifer test analysis software (Geraghty & Miller, 1994). Analysis results are presented in Appendix D and discussed in Section 3.3.

2.6 EQUIPMENT DECONTAMINATION PROCEDURES

All downhole soil sampling tools (e.g., stainless steel Geoprobe® drive-shoe and sampling barrel) were cleaned prior to collection of each sample with a clean water/phosphate-free detergent mix followed by a clean water rinse. Decontaminated tools also were used for soil gas sampling.

A new, disposable Teflon® bailer was used to collect the groundwater sample from each well. The water level indicator probe and purge pump were decontaminated prior to each use with a clean water/phosphate-free detergent mix followed by a distilled water rinse.

2.7 INVESTIGATION-DERIVED WASTES (IDW)

Soil cuttings and unused soil samples were moved to an approved on-base storage area for later disposal by the Base. Purge water was discharged to the influent of the groundwater treatment system at the Base Exchange Service Station on 7th Street.

2.8 ANALYTICAL DATA QUALITY ASSESSMENT

2.8.1 Introduction

A Parsons ES electronic Level III validation was performed on the March 1998 analytical results obtained from the fixed-base laboratories. The validation included internal data checks and application of data qualifiers to the analytical results based on adherence to method protocols and project-specific control limits. Method protocols reviewed included:

- Analytical holding times,
- Method blanks,
- Trip blanks,
- Surrogate spikes,
- Matrix spikes/matrix spike duplicates (MS/MSDs),
- Laboratory control samples (LCSs), and

• Sample temperatures during shipping and storage.

Data qualifiers were applied to analytical results during the data validation process. All data were validated using method applicable guidelines and in accordance with the National Functional Guidelines for Organic Data Review (USEPA, 1994a) and the National Functional Guidelines for Inorganic Data Review (USEPA, 1994b). The following definitions provide explanations of the USEPA (1994a and 1994b) qualifiers assigned to analytical results during data validation. The data qualifiers described were applied to both inorganic and organic results.

- U The analyte was not present above the reported sample quantitation limit (SQL).
- J1 The analyte is qualified as an estimated value solely because it is greater than the MDL and less than the practical quantitation limit (PQL), indicating no laboratory quality issues.

2.8.2 Data Quality

Data quality for each QC parameter where exceptions were noted during the validation is summarized in this section. Only results that exceeded quality assurance (QA)/QC criteria are presented. All frequency requirements for collection of field QA/QC samples (MS/MSDs and blanks) were met. The frequency requirements for laboratory specific method QA/QC also were met.

Samples were collected and analyzed as specified in the methods. All samples are representative of the site and comparable with the results of previous and future investigations (when used in accordance with the validation qualifiers).

All sample results qualified as "U" or "J1" and used in accordance with the data validation qualifiers applied are usable for the intended purposes. Results qualified as "J1" represent an association to non-compliant QC criteria which has caused the reported concentration to be estimated. Project objectives do not exclude the use of estimated concentrations, and therefore the data value is usable for project purposes.

In summary, accuracy and precision were in control. All method specific criteria were in control.

SECTION 3

PHYSICAL CHARACTERISTICS OF THE STUDY AREA

This section describes the physical characteristics of the Military Gas Station and adjacent environs at Eglin AFB, as determined from data collected during previous site investigations (EA, 1993a; EA, 1993b; EA, 1994a; EA, 1994b) and by Parsons ES in March 1998 as part of the risk-based investigation. A summary of site characterization activities completed by Parsons ES to supplement existing data is presented in Section 2 of this CAP.

3.1 PHYSIOGRAPHY

Eglin AFB is located in the East Gulf Coastal Plain Physiographic Province. This province is characterized by relatively low topographic relief and a gradual slope toward the Gulf of Mexico. White sand beaches and sand ridges typically border the coastline, while flatlands and swamps extend 10 to 15 miles inland.

3.2 REGIONAL GEOLOGY AND HYDROGEOLOGY

Surficial deposits are characterized by unconsolidated sands, silts, and clayey sands of Pliocene to Recent age which extend to a depth of approximately 60 feet below ground surface (bgs). The Surficial Aquifer System contains the undifferentiated Quaternary sediments and the Citronelle Formation. The typical lithology of the surficial aquifer is primarily fine to coarse quartz sand, with clay, silt and gravel, and clayey sand and sandy clay lenses. Limonite-cemented zones, shell beds, and carbonates are also common. The thickness of the surficial aquifer varies from 40 to 100 feet and the elevation of the water table varies from 0 to 30 feet above mean sea level (msl). The water within the Surficial Aquifer System is generally unconfined; however, beds of low permeability may cause semi-confined or locally confined conditions in its deeper parts. Water table elevations and horizontal gradients generally reflect contours of the land surface.

Below the surficial aquifer is the Pensacola Clay, a relatively impermeable unit separating the surficial aquifer from the Floridan Aquifer. The Floridan Aquifer System contains the Bruce Creek Limestone, St. Marks Formation, Chattahoochee Formation, Suwannee Limestone, Marianna Limestone and Ocala Limestone. The typical lithology of the Floridan Aquifer System is vuggy, fossiliferous, microcrystalline to granular, argillaceous to sandy, porous limestone and dolomite. The limestone and dolomite may be interbedded with dolomitic sand, silt and clay beds. The Ocala Limestone forms one of the most permeable zones within the Floridan Aquifer System. The extensive development of secondary porosity by dissolution and dolomitization has greatly increased the permeability of the unit. The thickness of the

Floridan Aquifer System varies from 900 to 1,000 feet and the elevation of the top of this unit varies from 270 to 320 feet below msl.

3.3 SITE GEOLOGY AND HYDROGEOLOGY

Site geology is characterized by approximately 60 feet of fine to coarse quartz sand with traces of silt and clay overlying the Pensacola Clay. Groundwater depth at the site is approximately 35 feet bgs. Groundwater surface elevations measured in March 1998 are summarized on Table 3.1 and depicted on Figure 3.1. Groundwater contour maps from previous investigations are presented in Appendix B. Groundwater flows toward the northeast at an average hydraulic gradient of approximately 0.0025 foot per foot (ft/ft) (Figure 3.1). Slug test data (Table 3.2) indicate that the hydraulic conductivity of the surficial deposits at the site ranges from approximately 15 to 56 feet per day (ft/day) with an average of approximately 34 ft/day. Literature values for the hydraulic conductivity of fine to coarse quartz sand range from 2.8 ft/day to 2,835 ft/day (Spitz and Moreno, 1996), so the slug test results are within the expected range for hydraulic conductivity at the site. Based on these data and an estimated effective porosity of 25 percent for sand (Spitz and Moreno, 1996), the average advective groundwater velocity at the site is 0.34 ft/day [124 feet per year (ft/yr)].

3.4 SITE TOPOGRAPHY AND SURFACE WATER HYDROLOGY

The Military Gas Station and the surrounding area has relatively flat topography, with ground elevations at the site at approximately 55 feet msl. Surface water hydrology around the site is dominated by the stormwater sewer system. The closest surface water body to the Military Gas Station is Weekly Pond, which is located approximately 1,200 yards to the southeast (Figure 1.2). The nearest downgradient surface water body is an inlet of Choctawhatchee Bay, which is located approximately 1,650 yards to the northeast (Figure 1.2).

3.5 CLIMATOLOGICAL CHARACTERISTICS

Eglin AFB has a humid, semitropical climate. Daily average temperature in the hottest summer months, July and August, may range from a low of 70 degrees Fahrenheit (°F) to a high of 88°F, with an average of approximately 82°F. Winters are mild, with occasional frost from November through February. During the coldest months, December and February, the temperature may be as low as 18°F or as high as 74°F, with the average around 50°F. Average annual rainfall is approximately 64 inches and ranges from 3.5 inches in October to almost 9 inches in July.

TNE

7 SE

TABLE 3.1

GROUNDWATER ELEVATIONS

Military Gas Station

Eglin AFB, Florida

	TOC a/	Depth	Groundwater
	Elevation	to Water	Elevation
Location	(ft msl) b/	(ft below TOC)	(ft msl)
EA-4	55.47	33.82	21.65
EA-5	55.24	33.81	21.43
EA-8	56.04	33.96	22.08
EA-10	54.72	33.42	21.30

Notes:

a/ TOC = top of casing

b/ ft msl = feet above mean sea level

S: \ES\Remed\BIOPLUME\EGLIN\FIGURES\98dn0590.dwg, 07/01/98 at

TABLE 3.2 SLUG TEST ANALYSIS RESULTS Military Gas Station Eglin AFB, Florida

			Hydraulic Conductivit	y
Well	Slug Test Number	ft/min ^{a/}	ft/day b/	cm/sec c/
EA-10	1	0.0104	14.95	0.0053
EA-10	2	0.0205	29.48	0.0104
EA-11	1	0.0392	56.39	0.0199

AVG	0.0233	33.60	0.0119

a/ ft/min = feet per minute

b/ ft/day = feet per day

c/ cm/sec = centimeters per second

SECTION 4

TIER 1 ANALYSIS AND IDENTIFICATION OF CHEMICALS OF POTENTIAL CONCERN

This section provides an overview of the regulatory requirements for a risk-based, tiered approach to identification of COPCs and reviews the preliminary conceptual site model (CSM) developed for the Military Gas Station in the work plan (Parsons ES, 1998) as a means of selecting appropriate regulatory screening criteria to identify COPCs in affected site media (i.e., chemicals present at concentrations that could pose a risk to human and/or ecological receptors exposed to the affected media). This section also presents a screening-level Tier 1 analysis used to select the COPCs that are the focus of this CAP. The COPCs for the Military Gas Station are identified in the Tier 1 analysis based on estimated risks to human health posed by maximum detected contaminant concentrations. Conservative land use and exposure assumptions are used in the Tier 1 screening analysis to ensure that the nature and extent of any COPCs that could pose a risk to receptors at or near the site are fully described (Section 5), and that these chemicals are fully evaluated in subsequent analyses through quantitative fate and transport and receptor exposure evaluations (Sections 6 and 7).

4.1 REGULATORY REVIEW OF THE TIER 1 SCREENING PROCESS

As an initial step in determining the necessity for remedial action at the Military Gas Station, representative concentrations of site contaminants are compared to the generic NFA-With-Conditions TCLs for soil and groundwater presented in Tables IV and V of the *Petroleum Contamination Site Cleanup Criteria* (FDEP, 1997). Contaminant soil concentrations must be below the Direct Exposure II and the leachability target levels presented in Table IV (based on applicable groundwater criteria specified in 62-770.680 (1)(c), FAC). Concentrations of COPCs in groundwater must be below background concentrations or less than levels presented in Table V. Maximum dissolved site contaminant concentrations also are compared to the Table IX Natural Attenuation Source Default Values. This comparison provides an initial assessment of the potential appropriateness of monitored natural attenuation as a remedial alternative.

Those analytes with site concentrations that exceed the appropriate TCLs for soil and groundwater are considered to be COPCs, and are retained for further analysis concerning the risk-reduction requirements for the site. The nature and extent of these COPCs are described more fully in Section 5. Qualitative and quantitative fate and transport analyses are presented in Section 6 to evaluate the migration and persistence of COPCs in affected media.

4.2 PRELIMINARY CONCEPTUAL SITE MODEL REVIEW

Figure 4.1 presents the preliminary conceptual site model (CSM) developed for the Military Gas Station. The CSM was developed using data collected during all relevant site investigations and is based on a review of potential receptors and feasible exposure scenarios. The purpose of developing a CSM is to guide the evaluation of available site information and to determine potential data gaps, including:

- Potential contaminant sources:
- Media affected by contaminant releases;
- Mechanisms of contaminant release (e.g., leaching and volatilization);
- Routes of possible receptor exposure (e.g., inhalation, ingestion, or dermal contact).
- · Potential human and ecological receptors; and
- Potential receptor exposure points based on conservative, reasonable land use assumptions.

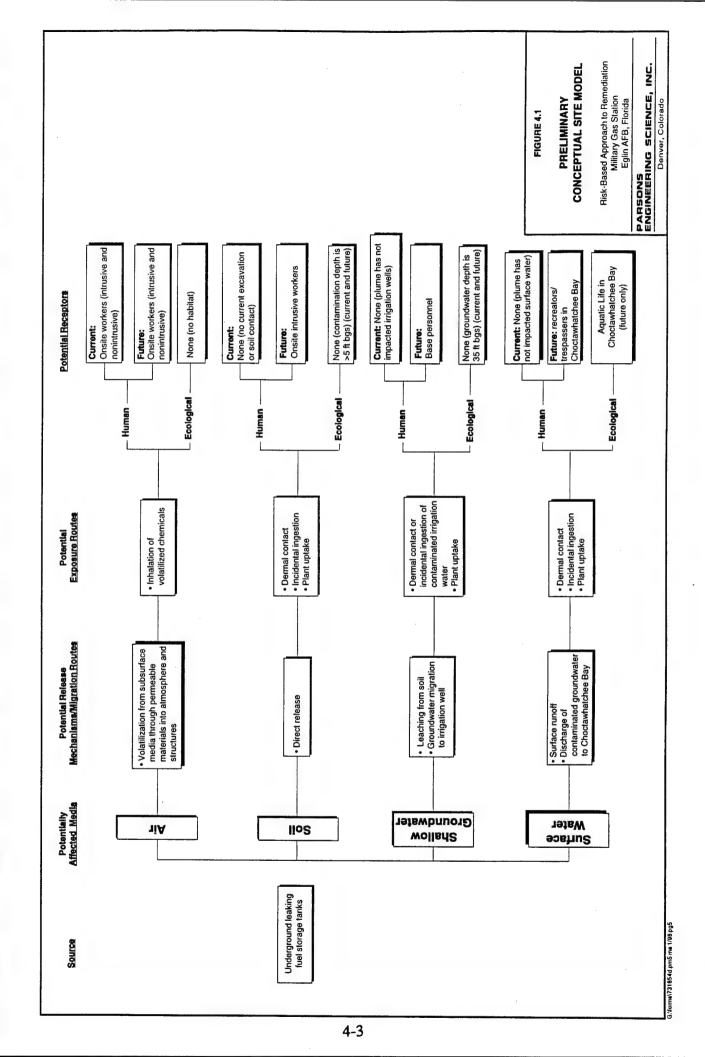
The CSM also was developed to provide an outline for addressing all media-specific, current and potential future exposure scenarios at the site. The CSM has been constructed to identify potentially completed receptor exposure pathways. For an exposure pathway to be completed, there must be a contaminant source, a release mechanism, a contaminant migration pathway, an exposure route, and a receptor. If any of these components is missing, the pathway is considered incomplete, and receptors are not at risk from exposure to site contaminants.

4.2.1 Contaminant Source Assessment

Contamination at the Military Gas Station is present as a result of past overfills of and/or leaks from the underground fuel storage tanks at the site. The USTs and surrounding contaminated soil have been removed from the site. Current soil quality data, which will be discussed in detail in Section 5, indicate that the continuing source of groundwater contamination at the site has been substantially reduced. Mobile light, non-aqueous phase liquid (LNAPL) (free product) has not been found at the site.

4.2.2 Land Use and Potential Receptors

The Military Gas Station is an active fueling station for military vehicles. This site is located within the industrial/commercial section of the main base at the intersection of Daytona Road and Okaloosa Avenue. Potential receptors include only onsite intrusive and non-intrusive industrial workers. Due to the developed, urbanized nature of the site and corresponding lack of habitat, there are no ecological receptors to be considered.



4.2.3 Exposure Pathways

An understanding of potential exposure pathways is important in determining how potential receptors could contact contaminated media and how that contact could result in the uptake of chemicals. An exposure pathway analysis reviews the contaminant sources, locations, and types of environmental releases in relation to population locations and activity patterns to determine the potentially significant pathways and routes of receptor exposure. A completed exposure pathway consists of four necessary elements:

- A source and mechanism of chemical release,
- An environmental transport medium,
- · A point of potential contact with a receptor, and
- A feasible route of exposure at the exposure point.

If one or more of these elements is missing, the pathway is incomplete and there is no exposure (and therefore, no risk).

Based on the industrial land use scenario and site-specific contaminated media information, the following human receptor exposure routes may potentially be completed and were evaluated during the data analysis process:

- Inhalation of volatilized contaminants by current or future onsite intrusive and nonintrusive workers;
- Dermal contact with or incidental ingestion of contaminated soil by future onsite intrusive workers (e.g., during future excavation activities);
- Dermal contact with or incidental ingestion of irrigation water by future Base personnel; and
- Dermal contact with or incidental ingestion of downgradient surface water by future recreators or trespassers at Choctawhatchee Bay.

Available information indicates that none of these pathways currently is completed. Therefore, only potential future exposures are assessed. Future receptors may be exposed to soil contaminants by dermal contact with and/or incidental ingestion of soil during intrusive activities (e.g., building foundation excavation, utility pipeline excavation). There are no completed exposure pathways for contaminated groundwater at the site. The depth to groundwater is approximately 35 ft bgs; therefore, groundwater will not be encountered during future excavation activities. No drinking water supply wells screened within the surficial aquifer are located within one-half mile of the site. Eight irrigation wells used to water base lawns are screened in the surficial aquifer within one-half mile of the site (EA, 1994) (Figure 1.2). However, as described in Section 6, the contaminant plume (which is currently wholly contained onsite) is not expected to migrate to these wells because the plume is shrinking.

Surface water may become contaminated by contact with contaminated soil or groundwater discharge to a surface water body. The surface water runoff pathway is not considered complete because the site is mostly paved and surface soils are not contaminated. Therefore, surface water runoff, which is channeled into the storm water drainage system, should not contact contaminated soils. The nearest surface water body downgradient from the site is an inlet of Choctawhatchee Bay located 1,650 yards to the northeast of the site. As described in Section 6, the dissolved contaminant plume is not expanding, and is wholly contained on-site. Therefore, site contamination is not expected to impact Choctawhatchee Bay. The site is expected to continue as a paved, urbanized environment; therefore, future risks to ecological receptors are unlikely.

4.3 TIER 1 SCREENING ANALYSIS

It is the intention of the Air Force to obtain FDEP approval for a corrective action for the site that will protect potential receptors from unacceptable exposures to site-related chemicals. To accomplish this objective, the COPCs that drive potential risks and impact the final remedial requirements at this site were identified.

FDEP (1997) Tier 1 TCLs are based on 1) analyte-specific toxicity data; 2) an exposure-pathway-specific cancer target risk limit of 10⁻⁶ (i.e., one additional cancer above the background rate in a population of one million) and a noncancer hazard quotient less than or equal to 1; and 3) conservative receptor exposure assumptions.

4.3.1 Tier 1 Screening Analysis for Soil

TCLs for direct exposure of industrial workers (Direct Exposure II) were selected as the appropriate set of Tier 1 screening values for soil at the Military Gas Station. The FDEP (1997) guidance provides industrial-scenario TCLs for petroleum constituents in soil that incorporate risks posed by the dermal contact, ingestion, and inhalation exposure pathways. Table 4.1 compares the maximum site concentrations for each compound measured in soil at the site during the 1998 risk-based sampling event to the Direct-Exposure II TCLs. The 1998 soil samples were collected from the locations containing the most elevated contaminant concentrations based on results from previous sampling events. Based on these comparisons, no analytes are identified as site COPCs in soil.

4.3.2 Tier 1 Screening Analysis for Groundwater

The Tier 1 groundwater TCLs presented by the FDEP (1997) and used in this CAP are based on the conservative assumption of unrestricted future use of groundwater (e.g., use as a drinking water source). Comparisons of the TCLs for unrestricted groundwater use to March 1998 data are presented in Table 4.2. Based on these comparisons, ethylbenzene, total xylenes, lead, and naphthalene are identified as the COPCs in site groundwater. Only the total xylenes concentration in source area well EA-4 exceeded its natural attenuation source default criterion. It should be noted that future use of groundwater as a drinking water source is not anticipated. Therefore, the Tier 1 TCLs are not realistic short-term cleanup levels for the site, and are used for preliminary screening purposes only.

TABLE 4.1 TIER 1 SCREENING SUMMARY FOR SOIL

Military Gas Station Eglin AFB, Florida

Analyte	Units	Maximum Concentration	Location (and Depth Interval in ft bgs) of Concentration	Direct Exposure ^{a/} II	Leachability b/
Benzene	mg/kg c/	0.0061 U ^{d/}	NA e/	1.50	0.007
Ethylbenzene	mg/kg	0.005 U	NA	240	0.4
Toluene	mg/kg	0.0061 U	NA	2,000	0.4
Xylenes (total)	mg/kg	0.012	V (26 - 28)	290	0.3
Methyl-tert-butyl ether	mg/kg	0.0061 U	NA	6,100	0.2
TRPH (C8-C40) ^f	mg/kg	6.5 J1 ^{g/}	K (15 - 17)	2,500	340
Acenaphthene	mg/kg	0.24 U	NA	22,000	4
Acenaphthylene	mg/kg	0.24 U	NA	11,000	22
Anthracene	mg/kg	0.024 U	NA	290,000	2000
Benzo(a)anthracene	mg/kg	0.074	V (26 - 28)	5.1	2.9
Benzo(a)pyrene	mg/kg	0.069	V (26 - 28)	0.5	7.8
Benzo(b)fluoranthene	mg/kg	0.11	V (26 - 28)	5	9.8
Benzo(g,h,i)perylene	mg/kg	0.063	V (26 - 28)	45,000	13,000
Benzo(k)fluoranthene	mg/kg	0.037	V (26 - 28)	52	25
Chrysene	mg/kg	0.11	V (26 - 28)	490	80
Dibenz(a,h)anthracene	mg/kg	0.024 U	NA	0.5	14
Fluoranthene	mg/kg	0.17	V (26 - 28)	45,000	550
Fluorene	mg/kg	0.015 J1	V (26 - 28)	24,000	87
Indeno(1,2,3-cd)pyrene	mg/kg	0.036 U	NA	5.2	28
Naphthalene	mg/kg	0.24 U	NA	8,600	1
Phenanthrene	mg/kg	0.11	V (26 - 28)	29,000	120
Pyrene	mg/kg	0.16	V (26 - 28)	40,000	570

Notes:

- a/ Direct Exposure II = FDEP TCLs based on No Further Action With Conditions.
- b/ Leachability = Based on FDEP Table V Groundwater TCLs.
- c/ mg/kg = Milligrams per kilogram.
- d/ U = The analyte was analyzed for and is not present above the reporting limit.
- e/ NA = Not applicable.
- f/ TRPH = Total Recoverable Petroleum Hydrocarbons.
- g/ J1 = The analyte was positively identified and has a concentration between the method detection limit and the reporting limit.

TABLE 4.2 TIER 1 SCREENING VALUES FOR GROUNDWATER

Military Gas Station Eglin AFB, Florida

Analyte	Units	Maximum Detection	Detection Location	Table V Target Cleanup Level ^a '	Table IX Natural Attenuation Source Default "
Benzene	μg/L c/	10 U ^d (0.28) ^e	NA ^{f/}	1	100
Ethylbenzene	μg/L	76	EA-4	30	300
Toluene	μg/L	10 U	NA	40	400
Xylenes (total)	μg/L	400	EA-4	20	200
TRPH (C8-C40) ^{g/}	mg/Lh/	4.4	EA-4	5	50
EDB i/	μg/L	0.020 U	NA	0.02	2
Lead	μg/L	19	EA-10	15	150
Acenaphthene	μg/L	1 U	NA	20	200
Acenaphthylene	μg/L	1 U	NA	210	2,100
Anthracene	μg/L	0.1 U	NA	2,100	21,000
Benzo(a)anthracene	μg/L	0.13 U	NA	0.2	20
Benzo(a)pyrene	μg/L	0.23 U (0.066)	NA	0.2	20
Benzo(b)fluoranthene	μg/L	0.18 U	NA	0.2	20
Benzo(g,h,i)perylene	μg/L	0.2 U	NA	210	2,100
Benzo(k)fluoranthene	μg/L	0.17 U	NA	0.5	50
Chrysene	μg/L	0.2 U	NA	5	500
Dibenz(a,h)anthracene	μg/L	0.3 U (0.078)	NA	0.2	20
Fluoranthene	μg/L	0.2 U	NA	280	2,800
Fluorene	μg/L	0.085 J1 ^{j/}	EA-4	280	2,800
Indeno(1,2,3-cd)pyrene	μg/L	0.44 U (0.056)	NA	0.2	20
Naphthalene	μg/L	40	EA-4	20 g j	200
Phenanthrene	μg/L	0.2 U	NA	210	2,100
Pyrene	μg/L	0.2 U	NA	210	2,100

Notes:

- a/ Table V Target Cleanup Level = FDEP TCLs for Groundwater
- b/ Table IX Natural Attenuation Source Default = FDEP TCLs for Natural Attenuation.
- c/ μ g/L = micrograms per liter.
- d/ U = The analyte was analyzed for and is not present above the reporting limit.
- e/ When the reporting limit exceeds the target cleanup level and the chemical is not detected (U), then the method detection limit (MDL) is shown in parentheses. The analyte concentration is below the MDL.
- f/ NA = Not applicable.
- g/ TRPH = Total Recoverable Petroleum Hydrocarbons.
- h/ mg/L = milligrams per liter.
- i/ EDB = ethylene dibromide.
- j/ J1 = The analyte was positively identified and has a concentration between the method detection limit and the reporting limit.

4.3.3 Tier 1 Screening Analysis for Soil Gas

FDEP (1997) guidance does not provide RBSLs for screening soil gas concentrations or for directly screening ambient air values. FDEP guidance accounts for the potential for volatilization of contaminants from soils into ambient air in the calculation of the Tier 1 TCLs for direct contact with soil. There was no COPC detected above Tier 1 TCLs in soil, indicating that exposure via volatilization from soil into ambient air will not present a risk. The Tier 1 TCLs do not account for the presence of the pavement cover over much of the site, which would act to further minimize the potential for exposure via the inhalation pathway. Currently, there are no enclosed structures at the site, and therefore no indoor receptors.

As a secondary means of assessing the potential for exposure via inhalation of volatiles, the soil gas sample collected in March 1998 was analyzed for BTEX, and maximum detections of each compound were compared to the chemical-specific Occupational Safety and Health Administration (OSHA) 8-hour time-weighted average Permissible Exposure Limits (PELs) (NIOSH, 1997) and time-weighted average threshold limit values (TWA-TLVs) [American Conference of Government Industrial Hygienists (ACGIH), 1998]. Table 4.3 presents the results of this comparison. No BTEX constituents were detected above the OSHA PELs or the TLVs, indicating that inhalation of volatilized contaminants does not currently, and will not in the future, pose a risk to potential receptors.

4.3.4 Summary of Site COPCs

Based on comparisons of the maximum soil, groundwater, and soil gas concentrations to FDEP (1997) TCLs and OSHA PELs (NIOSH, 1997), dissolved ethylbenzene, dissolved xylenes, dissolved lead, and dissolved naphthalene are identified as COPCs for the Military Gas Station.

TABLE 4.3

TIER 1 SCREENING VALUES FOR SOIL GAS

Military Gas Station Eglin AFB, Florida

Analyte	Maximum Detection (ppmv ^{a/})	OSHA PEL ^{b/} (ppmv)	TWA-TLV ^{c'} (ppmv)
Benzene	< 0.020	1	0.5
Ethylbenzene	<0.020	200	100
Toluene	0.026	100	50
Xylenes (total)	0.075	100	100
TPH ^{d/}	1.5	Not available	Not available
C2-C4 Hydrocarbons e/	<0.20	Not available	Not available

Notes:

- a/ ppmv = parts per million, volume per volume.
- b/ OSHA PEL = Occupational Safety and Health Administration (NIOSH, 1997) 8-hour time-weighted average permissible exposure limit.
- c/ TLV = Time-weighted average/threshold limit value recommended by the American Conference of Government Industrial Hygienists (ACGIH), 1998.
- d/ TPH = Total Petroleum Hydrocarbons (C5+ hydrocarbons referenced to gasoline).
- e/ Referenced to gasoline.

SECTION 5

ANALYTICAL DATA SUMMARY AND MAGNITUDE AND EXTENT OF CHEMICALS OF POTENTIAL CONCERN

5.1 OVERVIEW

This section presents analytical results from the March 1998 field sampling event in tabular form, and summarizes the magnitude and extent of COPC contamination in sampled media at the Military Gas Station. Discussion in this section is primarily limited to those chemicals that were identified as COPCs based on the Tier 1 screening analysis presented in Section 4 (i.c., dissolved ethylbenzene, xylenes, lead, and naphthalene).

5.2 SOIL SAMPLING RESULTS

Soil sampling was performed as part of the recent risk-based investigation; analytical results are summarized in Table 5.1. Soil borings were advanced in areas of elevated fuel contamination (based on previous investigations) to determine worst case hydrocarbon concentrations in soil, in addition to facilitating evaluation of how concentrations have changed over time. As shown in Table 4.1, no analyte was detected above the Direct Exposure II levels. Available soil analytical data indicate that there are no exceedences of Tier I TCLs and, therefore, no soil COPCs.

5.3 GROUNDWATER SAMPLING RESULTS

Groundwater sampling was performed as part of the recent risk-based investigation; analytical results are summarized in Table 5.2. Sample locations were selected based on the results of previous investigations. The sampling objective was to determine the areal extent and magnitude of fuel hydrocarbon and lead concentrations in groundwater. As shown in Table 4.2, ethylbenzene, xylenes, lead, and naphthalene in groundwater exceeded their respective Tier I TCLs.

During the March 1998 field effort, ethylbenzene was detected above the FDEP TCL of 30 μ g/L at source area monitoring well EA-4 (76 μ g/L). Xylenes also were detected above the FDEP TCL of 20 μ g/L at EA-4 (400 μ g/L). Lead was detected above the FDEP TCL of 15 μ g/L at EA-4 (17 μ g/L) and at EA-10 (19 μ g/L). It should be noted, however, that only dissolved lead concentrations exceeded the TCL; total lead concentrations were below the TCL. Naphthalene was detected above the FDEP TCL of 20 μ g/L at EA-4 (40 μ g/L). The distribution of ethylbenzene, xylenes, lead, and naphthalene in groundwater is presented on Figure 5.1.

TABLE 5.1 SUMMARY OF SOIL ANALYTICAL DATA

Military Gas Station Eglin AFB, Florida

		Sample Locations, Depth Intervals (ft bgs), and Sampling Dates					
		В	V	v	K	К	
		25 - 27	8 - 10	26 - 28	8 - 10	15 - 17	
Analyte	Units	3/29/98	3/29/98	3/29/98	3/30/98	3/30/98	
Benzene	mg/kg a/	0.0061 U b/	NA c/	0.0061 U	NA	0.0055 U	
Ethylbenzene	mg/kg	0.0024 U	NA	0.005 U	NA	0.0022 U	
Toluene	mg/kg	0.0061 U	ŇA	0.0061 U	NA	0.0055 U	
Xylenes (total)	mg/kg	0.0061 U	NA	0.012	NA	0.0055 U	
Methyl-tert-butyl ether	mg/kg	0.0061 U	NA	0.0061 U	NA	0.0055 U	
TRPH (C8-C40) ^{d/}	mg/kg	4.4 J1 ^{e/}	NA	4.7 J1	NA	6.5 J1	
Acenaphthene	mg/kg	0.24 U	NA	0.24 U	NA	0.22 U	
Acenaphthylene	mg/kg	0.24 U	NA	0.24 U	NA	0.22 U	
Anthracene	mg/kg	0.024 U	NA	0.024 U	NA	0.022 U	
Benzo(a)anthracene	mg/kg	0.024 U	NA	0.074	NA	0.022 U	
Benzo(a)pyrene	mg/kg	0.018 U	NA	0.069	NA	0.016 U	
Benzo(b)fluoranthene	mg/kg	0.015 U	NA	0.11	NA	0.013 U	
Benzo(g,h,i)perylene	mg/kg	0.061 U	NA	0.063	NA	0.055 U	
Benzo(k)fluoranthene	mg/kg	0.013 U	NA	0.037	NA	0.012 U	
Chrysene	mg/kg	0.049 U	NA	0.11	NA	0.044 U	
Dibenz(a,h)anthracene	mg/kg	0.024 U	NA	0.024 U	NA	0.022 U	
Fluoranthene	mg/kg	0.049 U	NA	0.17	NA	0.044 U	
Fluorene	mg/kg	0.049 U	NA	0.015 J1	NA	0.044 U	
Indeno(1,2,3-cd)pyrene	mg/kg	0.036 U	NA	0.036 U	NA	0.033 U	
Naphthalene	mg/kg	0.24 U	NA	0.24 U	NA	0.22 U	
Phenanthrene	mg/kg	0.049 U	NA	0.11	NA	0.044 U	
Pyrene	mg/kg	0.049 U	NA	0.16	NA	0.044 U	
TOC ^{f/}	mg/kg	NA	2000 U	NA	2000 U	NA	

Notes

a/ mg/kg = Milligrams per kilogram.

b/ U = The analyte was analyzed for and is not present above the reporting limit.

c/ NA = Not Analyzed.

d/ TRPH = Total Recoverable Petroleum Hydrocarbons.

e/ JI = The analyte was positively identified and has a concentration between the method detection limit and the reporting limit.

f/ TOC = Total organic carbon.

TABLE 5.2 SUMMARY OF GROUNDWATER ANALYTICAL DATA Military Gas Station

Eglin AFB, Florida

		Sample Locations and Dates					
Analyte	Units	EA-4 3/31/98	EA-40 3/31/98 (Duplicate)	EA-5 3/27/98	EA-8 3/27/98	EA-10 3/27/98	
Benzene	μg/L ^{a/}	10 U ^{b/}	10 U	2 U	2 U	2 U	
		76	74	1.4 J1 ^{c/}	0.099 J1	2.3	
Ethylbenzene Toluene	μg/L μg/L	10 U	10 U	2 U	2 U	2.3 2 U	
Xylenes (total)	μg/L μg/L	400	380	6.7	4.7	11	
	7.8						
EDB ^d	μg/L	0.02 U	0.02 U	0.02 U	0.02 U	0.02 U	
TRPH (C8-C40) e/	mg/L ^{f/}	3.1	4.4	NA ^{g/}	NA	0.81	
Total Lead	μg/L	15	15	NA	NA	11	
Dissolved Lead	μg/L	17	17	NA	NA	19	
Acenaphthene	μg/L	0.98 U	1 U	NA	NA	1 U	
Acenaphthylene	μg/L	0.98 U	1 U	NA	NA	1 U	
Anthracene	μg/L	0.098 U	0.1 U	NA	NA	0.1 U	
Benzo(a)anthracene	μg/L	0.13 U	0.13 U	NA	NA	0.13 U	
Benzo(a)pyrene	μg/L	0.23 U	0.23 U	NA	NA	0.23 U	
Benzo(b)fluoranthene	μg/L	0.18 U	0.18 U	NA	NA	0.18 U	
Benzo(g,h,i)perylene	μg/L	0.2 U	0.2 U	NA	· NA	0.2 U	
Benzo(k)fluoranthene	μg/L	0.17 U	0.17 U	NA	NA	0.17 U	
Chrysene	μg/L	0.2 U	0.2 U	NA	NA	0.2 U	
Dibenz(a,h)anthracene	μg/L	0.29 U	0.31 U	NA	NA	0.3 U	
Fluoranthene	μg/L	0.2 U	0.2 U	NA	NA	0.2 U	
Fluorene	μg/L	0.056 J1	0.085 J1	NA	NA	0.2 U	
Indeno(1,2,3-cd)pyrene	μg/L	0.42 U	0.44 U	NA	NA	0.44 U	
Naphthalene	μg/L	40	27	NA	NA	0.7	
Phenanthrene	μg/L	0.2 U	0.2 U	NA	NA	0.2 U	
Pyrene	μg/L	0.2 U	0.2 U	NA	NA	0.2 U	

Notes:

a/ μ g/L = Micrograms per liter.

b/ U = The analyte was analyzed for and is not present above the reporting limit.

c/ Jl = The analyte was positively identified and has a concentration between the method detection limit and the reporting limit.

d/ EDB = Ethylene dibromide.

e/ TRPH = Total Recoverable Petroleum Hydrocarbons.

f/ mg/L = milligrams per liter.

g/ NA = Not analyzed.

S: \ES\Remed\BIOPLUME\EGLIN\FIGURES\98dn0592.dwg, 03/05/99 at 1

The distribution of xylenes presented on Figure 5.1 was used to estimate the volume of this compound present in the groundwater at the Military Gas Station. Calculations are presented in Appendix E. The estimated volume of xylenes in groundwater at the Military Gas Station is 134 mL (9.1 tablespoons [tbsp]).

5.4 SOIL GAS SAMPLING RESULTS

A soil gas sample was collected at the site to facilitate assessment of the potential risk to future workers at the site from inhalation of VOCs, and to determine whether or not sufficient O_2 is available in the soil gas to sustain aerobic fuel hydrocarbon biodegradation. The soil gas sample was collected at a depth of 3 feet bgs in the contaminant source area (Figure 2.1) and analyzed in the field for concentrations of O_2 and CO_2 . The sample also was submitted to Air Toxics, Ltd. of Folsom California for analysis of BTEX and TPH (referenced to gasoline and propane). Field and laboratory analytical results for the March 1998 soil gas sample are summarized in Table 5.3. Comparison of maximum soil gas BTEX concentrations to OSHA 8-hour time-weighted average PELs and TWA-TLVs (Table 4.3) indicates that hydrocarbons do not pose a potential inhalation risk to future intrusive or aboveground workers under current conditions.

It should be noted that the SVE system at the site was operating when the soil gas sample was collected, and may have affected soil gas contaminant concentrations. However, as noted in Section 4.3.1, the FDEP (1997) soil TCLs incorporate risks posed by the inhalation exposure pathway. The lack of TCL exceedences indicates that the remaining soil contamination does not pose an inhalation risk to potential receptors.

The detected O₂ concentration (14.8 percent) indicates that sufficient O₂ is present to sustain aerobic biodegradation of fuel residuals. The SVE system is creating an influx of oxygen from uncontaminated soils surrounding the site. This oxygen is promoting biodegradation of residual hydrocarbons. CO₂ is being produced during the microbially-mediated aerobic biodegradation of fuel hydrocarbons. The monitored CO₂ concentration of 3.2 percent is consistent with the ongoing biodegradation of fuel residuals in site soils.

TABLE 5.3 SUMMARY OF SOIL GAS ANALYTICAL DATA Military Gas Station

Eglin AFB, Florida

	S	Sample Locations, Dates, and	Units	
		Mil SG1 31-Mar-98		
Analyte	ppmv ^{a/}	mg/L ^{b/}	percent	
Benzene	ND ^{c/}	ND	NA ^d	
Toluene	0.026	0.099	NA	
Ethylbenzene	ND	ND	NA	
Xylenes (total)	0.075	0.330	NA	
TPH (C5+ Hydrocarbons) e/	1.5	6.2	NA	
C2-C4 Hydrocarbons	ND	ND	NA	
Oxygen	NA	NA	14.8	
Carbon Dioxide	NA	NA	3.2	

Notes:

- a/ ppmv = parts per million, volume per volume.
- b/ mg/L = milligrams per liter.
- c/ ND = not detected.
- d/ NA = not applicable.
- e/ TPH = total petroleum hydrocarbons.

SECTION 6

CHEMICAL FATE ASSESSMENT

6.1 INTRODUCTION

Biodegradation of dissolved fuel constituents and the future migration and persistence of the dissolved COPCs identified in Section 4 are assessed in this section to support development of a long-term monitoring (LTM) plan that can be used to ensure that downgradient receptors will not be impacted by the dissolved COPCs.

As used throughout this report, the term "remediation by natural attenuation" (RNA) refers to a subsurface contaminant remediation strategy that relies on natural physical, chemical, and biological mechanisms to control exposure of potential receptors to concentrations of contaminants in soils and groundwater that exceed regulatory levels. These mechanisms include the processes of advection, hydrodynamic dispersion, dilution from recharge, sorption, volatilization, and biodegradation, which facilitate RNA of a variety of anthropogenic chemicals.

This section summarizes and interprets specific site characterization data relevant to documenting the effectiveness of RNA at minimizing dissolved COPC migration and reducing COPC concentration, mass, and toxicity over time.

6.2 OPERATIVE MECHANISMS OF CONTAMINANT ATTENUATION

Understanding the fate of COPC in environmental media is critical to evaluating and predicting contaminant distribution patterns. There are several physical, chemical, and biological processes that influence how a chemical behaves in soil and groundwater.

Nondestructive attenuation processes can be described as those physical and chemical processes that may prohibit significant contaminant migration but will not result in a permanent reduction in contaminant mass. Examples of nondestructive attenuation processes include volatilization, sorption, dilution from recharge, advection, and hydrodynamic dispersion. These processes must be evaluated when determining whether some type of remediation is warranted because chemical contamination poses or has the potential to pose a risk to human or ecological receptors. If contamination cannot reach a potential receptor exposure point, the contamination poses no risk.

In comparison to nondestructive chemical attenuation processes, destructive chemical attenuation processes result in the permanent removal of contaminant mass from the environment. Documenting and distinguishing the effects of destructive attenuation processes, such as biodegradation, from nondestructive attenuation processes is critical

to evaluating the potential for RNA to bring about a reduction in contaminant mass over time. The effectiveness of destructive attenuation processes at reducing contaminant mass at a site depends on how susceptible the chemical is to biodegradation and whether the site is characterized by physical, chemical, and biological conditions favorable to such processes.

Numerous laboratory and field studies have shown that hydrocarbon-degrading bacteria can participate in the degradation of many of the chemical components of different types of fuels (e.g., gasoline) under both aerobic and anaerobic conditions. Biodegradation of fuel hydrocarbons will occur when an indigenous population of hydrocarbon-degrading microorganisms is present in the soil and groundwater, and sufficient concentrations of electron acceptors and nutrients, including fuel hydrocarbons, are available to these organisms. Soils and groundwater with a history of exposure to fuel hydrocarbon compounds, such as at the Military Gas Station, generally contain microbial populations capable of facilitating biodegradation reactions (Wiedemeier *et al.*, 1995). The chemical basis for the biodegradation of BTEX is described in more detail in Section 6.4, where geochemical data relevant to documenting biodegradation at the field scale at the Military Gas Station are presented.

6.3 EVIDENCE OF CONTAMINANT REDUCTION OVER TIME

The first step in determining whether contaminant concentrations are being reduced in soils and groundwater at the Military Gas Station was to compare contaminant concentrations at selected sampling locations over time. The purpose of this comparison was to assess the evidence of field-scale contaminant mass loss. Decreases in the magnitude of contaminant concentrations at a site over time that cannot be explained by physical processes (e.g., source removal actions such as SVE, air sparging, mass transport in groundwater) may be the first indication that contaminants are biodegrading at the site.

6.3.1 VOC Concentration Trends in Soil

There are no historical soil contamination laboratory data to compare to the March 1998 data to demonstrate the effects of biodegradation and SVE. However, field screening data are available to indicate the changes in volatile soil contaminant concentrations at the site. In 1994, soil borings were advanced at locations B, V, and K. Organic vapor analyzer (OVA) readings were recorded on the 1994 boring logs. These data are compared in Table 6.1 to the field screening results for the 1998 borings that were advanced at adjacent locations. The data indicate that soil contaminant concentrations have been substantially reduced since 1994 due to the effects of biodegradation and SVE.

6.3.2 COPC Concentration Trends in Groundwater

COPC concentrations measured at select monitoring wells from September 1992 to March 1998 are summarized in Table 6.2. Ethylbenzene, total xylenes, and naphthalene concentrations over time are plotted on Figures 6.1 and 6.2. Figure 6.1 is a plot of the concentrations of the contaminants over time at source area well EA-4. The plot shows an overall decrease in the concentrations of the contaminants since

TABLE 6.1 COMPARISON OF 1994 AND 1998 SOIL HEADSPACE READINGS Military Gas Station

Eglin AFB, Florida

2511112) 1101144						
Boring ID	Sample Depth	1994 Headspace Result (ppmv ^{a/})	1998 Headspace Result (ppmv)			
В	25-27 ft bgs b/	93	60			
V	26-28 ft bgs	>1,000	20			
K	15-17 ft bgs	88	20			

^{a/} ppmv = parts per million, volume per volume.

b/ ft bgs = feet below ground surface.

TABLE 6.2 SUMMARY OF HISTORICAL DISSOLVED COPC CONCENTRATIONS

Military Gas Station Eglin AFB, Florida

WELL	DATE	ETHYLBENZENE (μg/L) a/	TOTAL XYLENES (μg/L)	NAPHTHALENE (μg/L)	LEAD (μg/L)
EA-4	Sep-92	940	4900	190	13
	Sep-93	510	2380	170	NA b/
	Jun-94	440	2100	75	NA
	Dec-95	. 310	1820	300	NA
	Aug-96	480	2060	150	NA
	Nov-96	170	1010	11	NA
	Feb-97	240	1590	120	NA
	Mar-98	76	400	40	17

WELL	DATE	ETHYLBENZENE (µg/L)	TOTAL XYLENES (µg/L)	NAPHTHALENE (μg/L)	LEAD (μg/L)
EA-10	Jul-93	210	1000	62	3.9
	Jun-94	390	1440	96	NA
	Dec-95	190	915	190	NA
	Mar-98	2.3	11	0.7	19

WELL	DATE	ETHYLBENZENE (μg/L)	TOTAL XYLENES (µg/L)	NAPHTHALENE (μg/L)	LEAD (μg/L)
EA-1	Sep-92	< 0.9	< 0.9	< 1.4	< 5
	Sep-93	140	543.6	51	NA
	Jun-94	160	320	8.7	NA
	Nov-96	< 1	< 1	< 1	NA
	Feb-97	< 1	< 1	< 1	NA

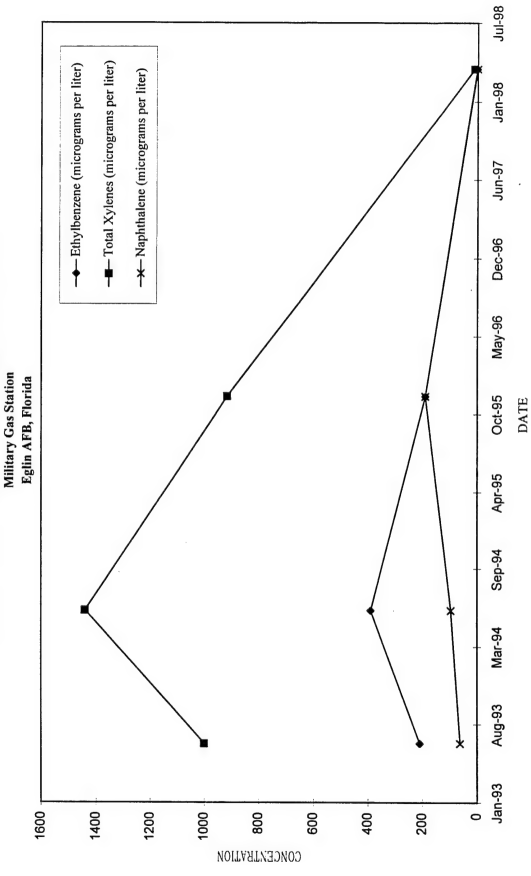
a/ μ g/L = micrograms per liter.

b/ NA = not analyzed.

• Expon. (Total Xylenes (micrograms per liter)) -Expon. (Ethylbenzene (micrograms per liter)) - Expon. (Naphthalene (micrograms per liter)) - Total Xylenes (micrograms per liter) - Ethylbenzene (micrograms per liter) Mar-97 - Naphthalene (micrograms per liter) COPC CONCENTRATIONS VERSUS TIME AT WELL EA-4 Oct-95 Military Gas Station Eglin AFB, Florida FIGURE 6.1 DATE $y = 7E + 16e^{-0.0009x}$ Jun-94 $y = 2E + 17e^{-0.001x}$ Jan-93 $y = 6E + 12e^{-0.0007x}$ Sep-91 5000 4500 4000 3500 3000 2500 2000 1500 1000 500

CONCENTRATION

COPC CONCENTRATIONS VERSUS TIME AT WELL EA-10



1992. Ethylbenzene concentrations have decreased from 940 $\mu g/L$ in 1992 to 76 $\mu g/L$ in 1998. Xylene concentrations have decreased from 4,900 $\mu g/L$ in 1992 to 400 $\mu g/L$ in 1998. Naphthalene concentrations have decreased from 190 $\mu g/L$ in 1992 to 40 $\mu g/L$ in 1998.

Exponential best-fit, first-order trendlines interpolated over the data for well EA-4 indicate contaminant reduction rates ranging from 0.0007 day for naphthalene (half-life = 2.7 years) to 0.001 day for ethylbenzene (half-life = 1.9 years). As with a large number of biological processes, the change in solute concentrations over time can generally be described using a first-order rate constant. The estimated time for the COPCs to decrease to below TCLs was calculated using the COPC-specific equations shown on Figure 6.1. Calculations are presented in Appendix E. The results indicate that the ethylbenzene concentration will decrease below the FDEP TCL of 30 μ g/L in 2.5 years, the total xylenes concentration will be below the FDEP TCL of 20 μ g/L in 2.7 years, and the naphthalene concentration will be below the FDEP TCL of 20 μ g/L in 2.7 years. It should be noted that substantial decreases in COPC concentrations occurred prior to startup of the air sparging/SVE system in 1994 or 1995, most likely due to the effects of biodegradation and removal of the USTs in 1991. It should also be noted that well EA-4 is located upgradient from the air sparging well, and the observed contaminant reduction rate should not be influenced by the sparging.

Figure 6.2 is a plot of the concentrations of ethylbenzene, total xylenes, and naphthalene over time at well EA-10. The plot shows an overall decrease in the concentration of COPCs since 1993. Ethylbenzene concentrations have decreased from 210 μ g/L in 1993 to 2.3 μ g/L in 1998. Xylene concentrations have decreased from 1,000 μ g/L in 1993 to 11 μ g/L in 1998. Naphthalene concentrations have decreased from 62 μ g/L in 1993 to 0.7 μ g/L in 1998.

6.4 EVIDENCE OF CONTAMINANT BIODEGRADATION VIA MICROBIALLY MEDIATED REDOX REACTIONS

Fuel hydrocarbons are typically utilized as electron donors in biologically mediated redox reactions under a wide range of geochemical conditions. Therefore, analytical data on potential electron acceptors can be used as geochemical indicators of fuel hydrocarbon biodegradation (Wiedemeier et al., 1995). Reductions in the concentrations of oxidized chemical species that are used by microorganisms to facilitate the oxidation of fuel hydrocarbon compounds within contaminated media are an indication that contaminants are biodegrading. Alternately, an increase in the metabolic byproducts resulting from the reduction of electron acceptors can be used as an indicator of contaminant biodegradation. The availability of potential electron acceptors to participate in contaminant biodegradation reactions can be used to estimate the total contaminant mass that can be biodegraded over time at this site. information can be used to predict how much dissolved COPC mass can be removed from saturated soils and groundwater at the Military Gas Station as a result of natural processes.

6.4.1 Relevance of Redox Couples in Biodegradation

Microorganisms obtain energy to replenish enzymatic systems and to reproduce by oxidizing organic matter. Biodegradation of dissolved fuel hydrocarbons is the result of a series of redox reactions that maintain the charge balance within the natural environment. Microorganisms facilitate the degradation of these organic compounds by transferring electrons from the electron donor (i.e., fuel hydrocarbons and native organic carbon) to available electron acceptors. Electron acceptors are elements or compounds that occur in relatively oxidized states and can participate in redox reactions involving these available electron donors. Electron acceptors known to be present in saturated soil and groundwater at the Military Gas Station are oxygen, nitrate/nitrogen, sulfate, ferric iron, and carbon dioxide.

Microorganisms facilitate fuel hydrocarbon biodegradation to produce energy for their use. The amount of energy that can be released when a reaction occurs or is required to drive the reaction to completion is quantified by the free energy of the reaction (Stumm and Morgan, 1981). Microorganisms are able to utilize electron transport systems and chemiosmosis to combine energetically favorable and unfavorable reactions to produce energy for life processes (i.e., cell production and maintenance). Microorganisms will facilitate only those redox reactions that will yield energy. By coupling the oxidation of fuel hydrocarbon compounds, which requires energy, to the reduction of other compounds (e.g., oxygen, nitrate/nitrite, manganese, ferric iron, sulfate, and carbon dioxide), which yields energy, the overall reaction will yield energy. Detailed information on the redox reactions required to biodegrade dissolved COPCs is included in Table 6.3. The reader is encouraged to review this information to more fully understand the chemical basis of biodegradation.

Figure 6.3 illustrates the sequence of microbially mediated redox processes based on the amount of free energy released for microbial use. In general, reactions yielding more energy tend to take precedence over processes that yield less energy (Stumm and Morgan, 1981). As Figure 6.3 shows, oxygen reduction would be expected to occur in an aerobic environment with microorganisms capable of aerobic respiration because oxygen reduction yields significant energy. However, once the available oxygen is depleted and anaerobic conditions dominate the interior regions of the contaminant plume, anaerobic microorganisms can utilize other electron acceptors in the following order of preference: nitrate/nitrite, manganese, ferric iron, sulfate, and finally carbon dioxide. Each successive redox reaction provides less energy to the system, and each step down in redox energy yield would have to be paralleled by an ecological succession of microorganisms capable of facilitating the pertinent redox reactions.

The expected sequence of redox processes can be estimated by the oxidation/reduction potential (ORP) of the groundwater. The ORP measures the relative tendency of a solution or chemical reaction to accept or transfer electrons, and can be measured in the field. This measurement can be used as a crude indicator of which redox reactions may be operating at a site. High ORPs mean that the solution (or available redox couple) has a relatively high oxidizing potential.

Microorganisms can only facilitate the biodegradation (oxidation) of the fuel hydrocarbon compounds using redox couples that have a higher ORP than the

TABLE 6.3 COUPLED OXIDATION REACTIONS MILITARY GAS STATION EGLIN AFB, FLORIDA

Coupled Ethylbenzene Oxidation Reactions	ΔG°, (kcal/mole Ethyl-benzene)	ΔG°, (kJ/mole Ethyl- benzene)	Stoichiometric Mass Ratio of Electron Acceptor to Compound
$10.5 O_2 + C_6 H_5 C_2 H_5 \Rightarrow 8 CO_{2,g} + 5 H_2 O$ Ethylbenzene oxidation /aerobic respiration	-1066.13	-4461	3.17:1
$8.4NO_3 + 8.4H^+ + C_6H_5C_2H_5 \Rightarrow 8CO_{2,g} + 9.2H_2O + 4.2N_{2,g}$ Ethylbenzene oxidation / denitrification	-1080.76	-4522	4.92:1
$42 H^{+} + 21 MnO_{2} + C_{6}H_{5}C_{2}H_{5} \Rightarrow 8 CO_{2}g + 21 Mn^{2+} + 26 H_{2}O$ Ethylbenzene oxidation / manganese reduction	-1066.27	-4461	17.24:1
$84H^{+} + 42Fe(OH)_{3,a} + C_{6}H_{5}C_{2}H_{5} \Rightarrow 8CO_{2} + 42Fe^{2+} + 110H_{2}O$ Ethylbenzene oxidation / iron reduction	-778.48	-3257	22:1 ^{a/}
$10.5H^+ + 5.25SO_4^2 + C_6H_5C_2H_5 \Rightarrow 8CO_{2g} + 5.25H_2S^o + 5H_2O$ Eth Ethylbenzene oxidation / sulfate reduction	-166.75	-697.7	4.75:1
$5.5H_2O + C_6H_3C_2H_3 \Rightarrow 2.75CO_{2,g} + 5.25CH_4$ Ethylbenzene oxidation / methanogenesis	-39.83	-166.7	0.79:1 ^{td}

Coupled Xylene Oxidation Reactions	ΔG° _r (kcal/mole <i>m</i> -xylene)	ΔG°_{r} (kJ/mole <i>m</i> -xylene)	Stoichiometric Mass Ratio of Electron Acceptor to Compound
$10.5 O_2 + C_6 H_4 (CH_3)_2 \Rightarrow 8 CO_{2g} + 5 H_2 O$ m-Xylene oxidation /aerobic respiration	-1063.25	-4448	3.17:1
$8.4NO_3 + 8.4H^+ + C_6H_4(CH_3)_2 \Rightarrow 8CO_{2,g} + 9.2H_2O + 4.2N_{2,g}$ m-Xylene oxidation / denitrification	-1077.81	-4509	4.92:1
$46 H^{+} + 22 \underline{MnO_2 + C_6H_4(CH_3)_2} \Rightarrow 8CO_{2g} + 22 \underline{Mn^2 + 28H_2O}$ $m\text{-Xylene oxidation / manganese reduction}$	-1063.39	-4449	11.39:1
$84H^{+} + 42Fe(OH)_{3,a} + C_{6}H_{4}(CH_{3})_{2} \Rightarrow 8CO_{2} + 42Fe^{2+} + 110H_{2}O$ $m-Xylene oxidation / iron reduction$	-775.61	-3245	22:1 ^{a/}
$10.5H^+ + 5.25SO_4^{2-} + C_6H_4(CH_3)_2 \Rightarrow 8CO_{2,g} + 5.25H_2S^o + 5H_2O$ m-Xylene oxidation / sulfate reduction	-163.87	-685.6	4.75:1
$5.5 H_2O + C_6H_4(CH_3)_2 \Rightarrow 2.75 CO_{2,g} + 5.25 CH_4$ m-Xylene oxidation / methanogenesis	-36.95	-154.6	0.79:1 ^ы

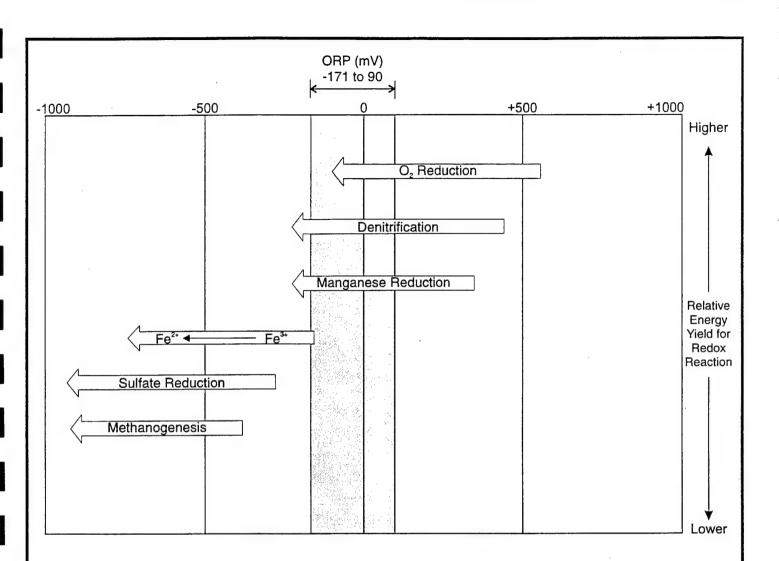
(Continued)

TABLE 6.3 (Continued) COUPLED OXIDATION REACTIONS MILITARY GAS STATION EGLIN AFB, FLORIDA

Coupled Naphthalene Oxidation Reactions	ΔG° _r (kcal/mole naphthalene)	ΔG°, (kJ/mole naphthalene)	Stoichiometric Mass Ratio of Electron Acceptor to Compound
$12O_2 + C_{10}H_8 \Rightarrow 10CO_2 + 4H_2O$ Naphthalene oxidation /aerobic respiration	-1217.40	-5094	3.00:1
$9.6NO_3^- + 9.6H^+ + C_{10}H_8 \Rightarrow 10CO_2 + 8.8H_2O + 4.8N_{2,g}$ Naphthalene oxidation / denitrification	-1234.04	-5163	4.65:1
$24MnO_2 + 48H^+ + C_{10}H_8 \Rightarrow 10CO_2 + 24Mn^2 + 28H_2O$ Naphthalene oxidation / manganese reduction	-1217.57	-5094	16.31:1
48Fe(OH) _{3,a} + 96H* + $C_{10}H_8 \Rightarrow 10CO_2 + 48Fe^{2+} + 124H_2O$ Naphthalene oxidation / iron reduction	-932.64	-3902	40.13:1
$6SO_4^{2c} + 12H^* + C_{10}H_8 \Rightarrow 10CO_2 + 6H_2S^o + 4H_2O$ Naphthalene oxidation / sulfate reduction	-196.98	-824.2	4.50:1
$8H_2O + C_{10}H_8 \Rightarrow 4CO_2 + 6CH_4$ Naphthalene oxidation / methanogenesis	-44.49	-186.1	0.75:1

a Mass of ferrous iron produced during microbial respiration.

^{b/} Mass of methane produced during microbial respiration.



Notes

ORP = Oxidation Reduction Potential

Range of ORP measured at the Military Gas Station

- These reactions would be expected to occur in sequence if the system is moving toward equilibrium.
- These redox processes occur in order of their energy-yielding potential (provided microorganisms are available to mediate a specific reaction).
 Reduction of a highly oxidized species decreases the ORP of the system.
- 3. The ORP of the system determines which electron acceptors are available for organic carbon oxidation.
- 4. Redox sequence is paralleled by an ecological succession of biological mediators.

FIGURE 6.3

SEQUENCE OF MICROBIALLY MEDIATED REDOX PROCESSES

Risk-Based Approach to Remediation Military Gas Station Eglin AFB, Florida

PARSONS ENGINEERING SCIENCE, INC.

Denver, Colorado

Adapted from Stumm and Morgan, 1981.

contaminants. This is why these electron acceptors can be used to oxidize the fuel hydrocarbon compounds. The reduction of highly oxidized species results in an overall decrease in the oxidizing potential of the groundwater. As shown in Figure 6.3, the reduction of oxygen and nitrate will reduce the oxidizing potential to levels at which ferric iron (Fe³⁺) reduction can occur. As each chemical species that can be used to oxidize the contaminants is exhausted, the microorganisms are forced to use other available electron acceptors with lower oxidizing capacity. When sufficiently low (negative) ORP levels have been developed as a result of these redox reactions, sulfate reduction, and methanogenesis can occur almost simultaneously (Stumm and Morgan, 1981).

ORP values measured in shallow groundwater at the Military Gas Station in March 1998 ranged from -171 to 90 millivolts (mV) (Table 6.4). Areas with the lowest ORP measurements generally coincided with the presence of fuel-contaminated groundwater, indicating that the progressive use of electron acceptors in the order shown on Figure 6.3 has caused the groundwater in the contaminated areas to become more reducing. These data imply that oxygen, nitrate, manganese, and ferric iron may be used to biodegrade fuel hydrocarbon contaminants at this site. However, many authors have noted that field ORP data alone cannot be used to reliably predict all of the electron acceptors that may be operating at a site, because the platinum electrode probes are not sensitive to some redox couples (e.g., sulfate/sulfide) (Stumm and Morgan, 1981; Godsey, 1994; Lovley et al., 1994). Analytical data on oxidized and reduced species are presented in the following subsections to verify which electron acceptors are actually being used to biodegrade the BTEX in saturated soil and groundwater at the Military Gas Station.

Throughout the following subsections, the distributions of geochemical parameters are examined by comparing background concentrations to BTEX plume core concentrations. Analytical data from upgradient well EA-8 and cross-gradient well EA-5 are used for background concentrations. Analytical data from EA-4 are used for BTEX plume core concentrations.

6.4.2 Dissolved Oxygen

Almost all types of fuel hydrocarbons can be biodegraded under aerobic conditions (Borden, 1994). Mineralization of fuel hydrocarbons to carbon dioxide and water under aerobic conditions involves the use of oxygen as a cosubstrate during the initial stages of metabolism, and as a terminal electron acceptor during the later stages of metabolism for energy production. The reduction of molecular oxygen during the oxidation of the fuel hydrocarbon compounds yields a significant amount of free energy that the microorganisms could utilize.

Dissolved oxygen (DO) concentrations were measured at groundwater sampling locations in March 1998. Table 6.4 presents the analytical results for DO by sampling location. The presence of the lowest observed DO concentration (1.2 mg/L) in the core of the dissolved hydrocarbon plume (EA-4) is an indication that biodegradation through aerobic respiration is occurring.

TABLE 6.4 SUMMARY OF GROUNDWATER GEOCHEMICAL DATA

	TABLE 6.4 SUMMARY OF GROUNDWATER GEOCHEMICAL DATA Military Gas Station Eglin AFB, Florida EA-4 EA-40 EA-5 EA-8 EA-10							
Parameter	Units	EA-4 31-Mar-98	EA-40 31-Mar-98	EA-5 27-Mar-98	EA-8 27-Mar-98	EA-10 27-Mar-98		
Ferrous Iron	mg/L a/	0.4	0.4	0.0	0.0	0.0		
Sulfate	mg/L	(67)	89	120	(109)	76		
Nitrate	mg/L	4.3	4.3	3.7	1.4	1.2		
Methane	mg/L	(1.7)	1.5	0.00054	(0.00038 J1 b)	0.0012		
Temperature	Deg C c/	25.0	NM ^{d/}	23.0	24.3	24.6		
pН	SU e/	6.21	NM	6.33	6.27	7.54		
Conductivity	μS/cm ^{f/}	0.130	NM	0.120	0.102	0.135		
Dissolved Oxygen	mg/L	1.2	NM	6.0	6.1	7.5		
ORP ^{g/}	mV ^{h/}	-171.4	NM	64.7	90.1	34.5		

Notes:

a/ mg/L = milligrams per liter.

b/ J1 = The analyte was positively identified and has a concentration between the method detection limit and the reporting limit.

- c/ Deg C = degrees Celcius.
- d/ NM = not measured.
- e/ SU = Standard Units.
- f/ μ S/cm = microsiemens per centimeter.
- g/ ORP = oxidation reduction potential.
- h/mV = millivolts.

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6.4.3 Nitrate

Once available DO concentrations are depleted through aerobic respiration, nitrate can be used as an electron acceptor by indigenous facultative anaerobes that mineralize fuel hydrocarbon compounds via either denitrification or nitrate reduction processes. Concentrations of nitrate (as nitrogen [N]) measured at the site in March 1998 are summarized in Table 6.4. Background nitrate (as N) concentrations measured in the [upgradient and cross-gradient wells ranged from 1.4 to 3.7 mg/L and averaged 2.6 mg/L. Conversely, nitrate (as N) concentrations measured in plume area wells ranged from 1.2 mg/L to 4.3 mg/L and averaged 2.8 mg/L. These data indicate that nitrate concentrations within the dissolved plume are not depleted relative to measured background concentrations. The results indicate that nitrate is not being used to oxidize fuel hydrocarbons in the anaerobic core of the dissolved plumes via denitrification or nitrate reduction.

6.4.4 Ferrous Iron

Although relatively little is known about the anaerobic metabolic pathways involving the reduction of ferric iron (Fe³⁺), this process has been shown to be a major metabolic pathway for some microorganisms (Lovley and Phillips, 1988; Chapelle, 1993). Elevated concentrations of ferrous iron (Fe²⁺) often are found in anaerobic, fuel-contaminated groundwater systems. Concentrations of dissolved ferrous iron once were attributed to the spontaneous and reversible reduction of ferric oxyhydroxides, which are thermodynamically unstable in the presence of organic compounds such as benzene. However, more recent studies suggest that the reduction of ferric iron cannot proceed at all without microbial mediation (Lovley and Phillips, 1988; Lovley et al., 1991; Chapelle, 1993). None of the common organic compounds found in low-temperature, neutral, reducing groundwater could reduce ferric oxyhydroxides to ferrous iron under sterile laboratory conditions (Lovley et al., 1991). This means that the reduction of ferric iron to ferrous iron requires mediation by microorganisms with the appropriate enzymatic capabilities.

To determine if ferric iron is being used as an electron acceptor for fuel biodegradation at the Military Gas Station, ferrous (reduced) iron concentrations were measured at groundwater sampling locations in March 1998. A low ferrous iron concentration (0.4 mg/L) was detected in the plume core well EA-4 (Table 6.4). Ferrous iron was not detected in groundwater from upgradient or cross-gradient wells. The occurrence of ferrous iron within the plume core indicates that ferric iron is acting as an electron acceptor at this location. In addition, the measured redox potentials of the groundwater at this site are within the range that would be expected for the ferric iron-reducing conditions implied by the observed ferrous iron distributions (Figure 6.3).

6.4.5 Sulfate

Sulfate also may be used as an electron acceptor during microbial degradation of fuel hydrocarbons under anaerobic conditions (Grbic-Galic, 1990). Sulfate can be reduced to sulfide during the oxidation of the fuel hydrocarbon compounds. The presence of decreased concentrations of sulfate (and increased concentrations of sulfide)

in the source area relative to background concentrations indicates that sulfate is participating in redox reactions at the site. To investigate the potential for sulfate reduction at the Military Gas Station, sulfa e concentrations were measured during the March 1998 groundwater sampling event.

Table 6.4 shows the analytical results for sulfate in groundwater at the Military Gas Station. In general, areas characterized by elevated concentrations of dissolved BTEX are depleted in sulfate concentrations relative to measured background concentrations. Background concentrations of sulfate at the site ranged from 109 mg/L at well EA-8 to 120 mg/L at EA-5. Sulfate concentrations measured at plume area wells exhibiting dissolved fuel contamination ranged from 67 mg/L at well EA-4 to 76 mg/L at well EA-10. This general depletion of sulfate within the contaminated areas indicates that this compound is acting as an electron acceptor during fuel biodegradation reactions.

The measured ORPs of the groundwater at this site are not within the range that would be expected for the sulfate-reducing conditions implied by the observed sulfate distributions. However, as described in Section 6.4.1, field ORP data alone cannot be used to reliably predict the electron acceptors that may be operating at a site.

6.4.6 Dissolved Methane

On the basis of free energy yield and the oxidizing potential of the site groundwater, the carbon dioxide/methane (CO₂/CH₄) redox couple also could be used to oxidize fuel hydrocarbon compounds to carbon dioxide and water once the groundwater is sufficiently reducing. To attain these reducing levels, other highly oxidizing chemical species such as oxygen, nitrate, ferric iron, and sulfate must first be reduced. This redox reaction is called methanogenesis or methane fermentation. Methanogenesis yields the least free energy to the system in comparison to other chemical species (Figure 6.3 and Table 6.3). The presence of methane in groundwater at elevated concentrations relative to background concentrations is a good indicator of methane fermentation.

Dissolved methane was measured at groundwater monitoring wells sampled during the March 1998 sampling event. Table 6.4 presents the analytical data for methane. Methane concentrations detected at the contaminant source area were elevated relative to background concentrations. The methane concentration measured at plume core area well EA-4 was 1.7 mg/L. In contrast, background concentrations were less than 0.00054 mg/L. The presence of elevated methane levels in groundwater at the Military Gas Station strongly indicates that biodegradation is occurring via methanogenesis.

6.4.7 pH

The pH of groundwater samples collected from groundwater monitoring points and monitoring wells in March 1998 was measured (Table 6.4). The pH of a solution is the negative logarithm of the hydrogen ion concentration [H⁺]. Groundwater pH values measured at the site were within the optimal range for fuel hydrocarbon-degrading microbes of 6 to 8.

6.4.8 Temperature

Groundwater temperature was measured at groundwater monitoring wells in March 1998 (Table 6.4). Temperature affects the types and growth rates of bacteria that can be supported in the groundwater environment, with higher temperatures generally resulting in higher growth rates. The temperature of groundwater samples collected from the shallow monitoring wells varied from 23 degrees Celsius (°C) to 25 °C. These relatively warm temperatures should promote microbial growth and may enhance rates of hydrocarbon biodegradation.

6.4.9 Theoretical Assimilative Capacity Estimates

The preceding discussions have been devoted to determining if fuel hydrocarbons are biodegrading in saturated soils and groundwater at the Military Gas Station. Analytical data on reduced and oxidized chemical species indicate that indigenous microorganisms are facilitating the oxidation of fuel hydrocarbons and the reduction of electron acceptors to generate free energy for cell maintenance and production. The question of how much contaminant mass can be biodegraded must be addressed to assess the full potential for long-term intrinsic bioremediation to minimize plume size and mass over time.

Mass balance relationships can be used to determine how much contaminant mass can be degraded by each of the redox reactions that the microorganisms might use to make free energy available for cell maintenance and production. The stoichiometric relationship between the contaminant and the electron acceptor can be used to estimate the expressed assimilative capacity of the groundwater. Once the redox reactions operating at the site have been defined, it is possible to estimate how much contaminant mass can be assimilated or oxidized by available electron acceptors.

Table 6.2 presents the coupled redox reactions that represent the biodegradation of the individual COPCs, including the stoichiometric mass ratio of electron acceptors needed to oxidize each compound. These stoichiometric mass ratios can be used to estimate the assimilative capacity of the groundwater at the Military Gas Station. For oxygen, nitrate, and sulfate, this is accomplished by first determining the initial (background) mass of each electron acceptor available in the groundwater. Data on these chemical species were collected at sampling locations upgradient and crossgradient from the dissolved plume. As groundwater slowly migrates into the source area, electron acceptors are brought into contact with hydrocarbon-degrading microorganisms and site contamination. The change in the electron acceptor mass from background sampling locations to sampling locations within the plume core is divided by the mass of electron acceptors required to mineralize the COPCs. For ferrous iron and methane, the highest observed concentration in the plume core wells is divided by the mass of electron acceptors required to mineralize the COPC. These numbers are summed to estimate the expressed intrinsic capacity of the groundwater to biodegrade each COPC.

Estimates of the background and plume core concentrations were used to calculate the expressed assimilative capacity of the groundwater system attributable to aerobic respiration, denitrification, and sulfate reduction (Table 6.5). The source area

ESTIMATED ASSIMILATIVE CAPACITY OF SATURATED SOIL AND GROUNDWATER TABLE 6.5

			Mass Ratio	Ethylbenzene
Electron Acceptor or	Background	Concentration in	of Electron Acceptor/	Assimilative
Metabolic Byproduct	Concentration	Core of Plume	Byproduct to COPCs b/	Capacity 6/
	(mg/L) a/	(mg/L)	(unitless)	(mg/L)
Oxygen	6.1	1.2	3.17	1.55
Nitrate	1.4	4.3	4.92	NA ^{d/}
Sulfate	109	29	4.75	8.84
Ferrous Iron	0.0	0.40	22	0.02
Methane	0.00038	1.7	0.79	2.15
			Total	12.56
		N	Max. 1998 Concentration (mg/L)	0.076

			Mass Ratio	Xylenes
Electron Acceptor or	Background	Concentration in	of Electron Acceptor/	Assimilative
Metabolic Byproduct	Concentration	Core of Plume	Byproduct to COPCs b/	Capacity 6/
	(mg/L) a/	(mg/L)	(unitless)	(mg/L)
Oxygen	6.1	1.2	3.17	1.55
Nitrate	1.4	4.3	4.92	NA
Sulfate	109	29	4.75	8.84
Ferrous Iron	0.0	0.40	22	0.02
Methane	0.00038	1.7	0.79	2.15
			Total	12.56
		N	Max. 1998 Concentration (mg/L)	0.400

(continued)

TABLE 6.5 (concluded) ESTIMATED ASSIMILATIVE CAPACITY OF SATURATED SOIL AND GROUNDWATER Military Gas Station Eglin AFB, Florida

0.040	Max. 1998 Concentration (mg/L)	W		
13.24	Total			
2.27	0.75	1.7	0.00038	Methane
0.01	40.13	0.40	0.0	Ferrous Iron
9.33	4.50		109	Sulfate
NA	4.65	4.3	1.4	Nitrate
1.63	3.00	1.2	6.1	Oxygen
(mg/L)	(unitless)	(mg/L)	(mg/L) a/	
Capacity c/	Byproduct to COPCs b/	Core of Plume	Concentration	Metabolic Byproduct
Assimilative	of Electron Acceptor/	Concentration in	Background	Electron Acceptor or
Naphthalene	Mass Ratio			

a/ mg/L = milligrams per liter.

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b/ Calculation based on the ratio of the total mass of electron acceptor required to oxidize a given average of the mass of contaminants of potential concern (ethylbenzene, total xylenes, and naphthalene).

c/ Assimilative capacity is the amount of contaminant that can be degraded by a certain method.

d/ NA = Not applicable.

13.2× = 33/ 0.04 + true no x Acc (40) concentrations of ferrous iron and methane are used to "back-calculate" the expressed assimilative capacity that is attributable to ferric iron reduction and methanogenesis. On the basis of these calculations, one pore volume of saturated soils and groundwater at the Military Gas Station has the capacity to oxidize an ethylbenzene and xylenes concentration of 12,560 μ g/L, and a naphthalene concentration of 13,250 μ g/L. As shown in Table 6.5, the assimilative capacity for each COPC is substantially higher than the maximum concentration of the COPC detected in groundwater at the Military Gas Station. The assimilative capacities should not change significantly if operation of the air sparging/SVE system is discontinued.

This estimate essentially represents an estimate of the reduction capability of one pore volume of groundwater at the Military Gas Station. The estimate identifies how much contaminant mass can be theoretically oxidized as one pore volume travels through the plume core. In reality, over 2 pore volumes are expected to move through the contaminated aquifer material in the source area each year based on the estimated groundwater velocity of 124 ft/yr.

A closed system containing 2 liters of water can be used to help visualize the physical meaning of assimilative capacity. Assume that the first liter contains no fuel hydrocarbons, but it contains fuel-degrading microorganisms and has an assimilative capacity of exactly "x" mg of fuel hydrocarbons. The second liter has no assimilative capacity; however, it contains fuel hydrocarbons. As long as these 2 liters of water are kept separate, biodegradation of fuel hydrocarbons will not occur. If these 2 liters are combined in a closed system, biodegradation will commence and continue until the fuel hydrocarbons or electron acceptors are depleted. If less than "x" mg of fuel hydrocarbons are in the second liter, all of the fuel hydrocarbons will eventually degrade given a sufficient time; likewise, if greater than "x" mg of fuel hydrocarbons would ultimately degrade.

This example shows that in a closed system, the measured expressed assimilative capacity eventually should be equivalent to the loss in contaminant mass; however, the groundwater beneath the site is an open system. Electron acceptors can continually enter the system from upgradient flow. Furthermore, contaminant mass can be added to the system through dissolution or leaching from LNAPL or contaminated soils. This means that the assimilative capacity is not fixed as it would be in a closed system, and therefore should not be quantitatively compared to concentrations of dissolved contaminants in the groundwater. Rather, the expressed assimilative capacity of groundwater is intended to serve as a qualitative tool. The fate of BTEX in groundwater is dependent on the relationship between the kinetics of biodegradation and the solute transport velocities (Chapelle, 1994).

6.5 COPC MIGRATION

The migration velocity of the groundwater COPCs ethylbenzene, xylenes, naphthalene, and lead, would be expected to be lower than the advective groundwater velocity computed in Section 3.3 (124 ft/yr) due to the effects of retardation. Retardation coefficients are calculated using the following formula:

where

 $K_d = (K_{oc})(f_{oc})$

 K_{∞} = Organic Carbon Partition Coefficient

 f_{∞} = Fraction Organic Carbon

 ρ_b = Soil Bulk Density of Aquifer Matrix

 n_e = Effective Porosity

Two soil samples collected at the Military Gas Station were analyzed for total organic carbon (TOC); however, as shown in Table 5.1, organic carbon was not detected. Similarly, organic carbon was not detected in three samples collected at the Seventh Street BX Service Station. If it is assumed that the TOC content of the shallow sand aquifer beneath the site is equal to one-half the method detection limit of 550 mg/kg, then the retardation coefficients of ethylbenzene, xylenes, and naphthalene would range from 1.75 to 2.04 (Table 6.6). The resulting migration velocities of ethylbenzene, xylenes, and naphthalene would be 66 ft/yr, 71 ft/yr, and 61 ft/yr, respectively based on an estimated advective groundwater velocity of 124 ft/yr. Lead typically adsorbs strongly to aquifer matrix materials such as manganese and iron oxide, organic carbon, or clay mineral surfaces, and is relatively immobile (Rose et al., 1979). Therefore, while lead resulting from gasoline contamination is not susceptible to degradation processes, it is generally restricted to the vicinity of the source area and does not pose a threat to offsite receptors.

Migration of dissolved contaminants can be assessed by comparing the estimated BTEX plume length over time. In June 1994, the dissolved BTEX plume extended approximately 450 feet downgradient from well EA-4 (see plume map in Appendix B). In contrast, the estimated plume length northeast of EA-4 in March 1998 was approximately 150 feet, indicating that the plume dimensions have been reduced considerably since 1994 due to the effects of biodegradation, air sparging, and source removal via SVE. In March 1998, Tier 1 TCLs were exceeded only in source area well EA-4; Tier 1 levels were not exceeded in groundwater from well EA-10, located approximately 120 feet downgradient from the source area.

TABLE 6.6 RETARDATION COEFFICIENTS OF COPCs

Military Gas Station Eglin AFB, Florida

		Average				ĺ
		Fraction		Bulk		
	K _{oc}	Organic		Density	Effective	
Compound	(L/kg ^{a/})	Carbon b/	Average ^{c3/}	(kg/L) ^{d/}	Porosity	Average
Ethylbenzene	468	0.000275	0.129	1.72	0.25	1.89
Xylenes	395	0.000275	0.109	1.72	0.25	1.75
Naphthalene	550	0.000275	0.151	1.72	0.25	2.04

- a/ L/kg = liters per kilogram.
- b/ Fraction organic carbon from accepted literature values.
- c/ Distribution coefficient
- d/ kg/L = kilograms per liter.

SECTION 7

TIER 2 ANALYSIS AND IDENTIFICATION OF FINAL CHEMICALS OF CONCERN

7.1 OBJECTIVE OF SITE-SPECIFIC EVALUATION

The Tier 1 analysis conducted in this CAP (Section 4) identified ethylbenzene, total xylenes, naphthalene, and dissolved lead as COPCs in groundwater. These analytes are evaluated in detail to better define/assess the potential adverse health effects they may cause in current or future human receptors.

The Tier 1 screening process is considered protective of human health because the Tier 1 risk-based screening criteria are based on conservative exposure assumptions. However, analytes identified as COPCs in Section 4 of this CAP (i.e., analytes with representative site concentrations exceeding Tier 1 TCLs) should not automatically be considered to be present at the Military Gas Station at levels that pose unacceptable threats to human health given the current and future exposure potential at this site. Rather, the exceedences of the screening criteria indicate that further evaluation using more site-specific exposure scenarios is warranted.

7.2 FINAL CONCEPTUAL SITE MODEL

The preliminary CSM presented in Section 4 was used to qualitatively identify potential human and ecological receptors that may be exposed to site-related contaminants, and to define the types of these potential exposures at or in the vicinity of the Military Gas Station (Figure 4.1). The preliminary CSM describes onsite release points, the affected physical media, the types of contaminant transport and fate mechanisms that may be involved at the site, each group of potentially exposed populations or receptors, and how each receptor group could come into contact with site-related contamination. This CSM was used to identify which of the exposure assumptions used to develop generic cleanup criteria most closely approximates site conditions. The exposure assumptions incorporated into the generic industrial TCLs (i.e., Table IV Direct-Exposure II TCLs) were identified as generally representative of the types of exposure that could occur at the Military Gas Station, but perhaps greatly overestimated the magnitude of exposure specific to current and expected future site conditions. For example, Tier 1 screening of groundwater assumed unrestricted future use of groundwater. Therefore, the target cleanup criteria presented in Table V (FDEP, 1997) which were developed assuming potable use of groundwater, were used in the Tier 1 screening. The preliminary CSM exposure pathways are reevaluated in this section using the Tier 2 chemical fate information presented in Section 6. It is important to emphasize that the purpose of using the preliminary CSM and the conservative, nonsite-specific TCLs to identify COPCs was to ensure that all subsequent assessment activities beyond the Tier 1 screening evaluation address the full range of contaminants that may present some risk to current of future receptors.

The revised CSM for the Military Gas Station, which is presented on Figure 7.1 and briefly reviewed in the following subsections, identifies only those receptors and exposure pathways that realistically may be involved in actual current or hypothetical future exposures. The outcome of the chemical fate assessment presented in Section 6 and the types of exposures likely to occur at this industrial site are reflected in this revised CSM. Justification for each site-specific exposure assumption is provided in subsequent discussions.

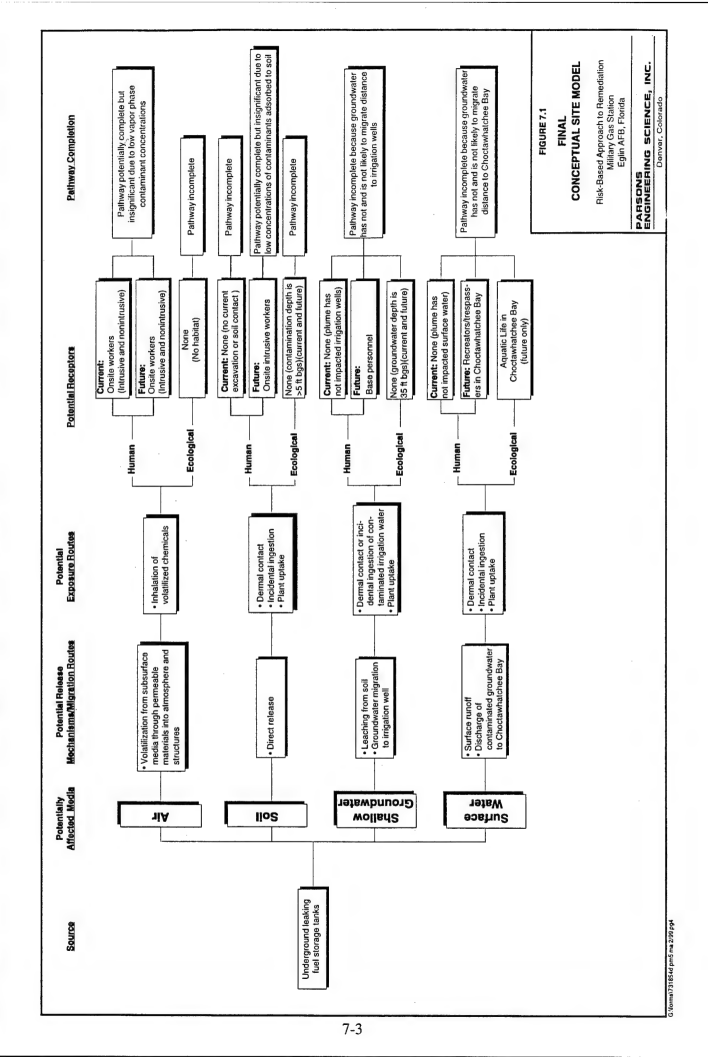
7.2.1 Sources, Affected Media, Release Mechanisms, and Contaminant Environmental Transport

The likelihood of release from a source, the nature of the contaminants involved, the affected environmental media, and the probable magnitude of their release all are included in the revised CSM (Figure 7.1). As described in the preliminary CSM (Figure 4.1), releases from the former gasoline USTs have contaminated site soil, soil gas, and groundwater with fuel hydrocarbons. The predominant ongoing release mechanism for groundwater COPCs is leaching from contaminated soils. The lack of mobile LNAPL (free product) detections at the site indicates that mobile LNAPL is not a significant, continuing source of groundwater contamination. Site data indicate that RNA and engineered remedial actions rates are also acting to limit migration of contaminants in concentrations above the TCLs; in fact, dissolved contaminant concentrations are being substantially reduced and the dissolved fuel hydrocarbon plume appears to be collapsing toward the source area. Therefore, future offsite migration of the contaminant plume is not anticipated. Therefore, dissolved contaminants will not impact Base irrigation wells or downgradient surface water bodies.

7.2.2 Potentially Exposed Receptors, Exposure Points, and Exposure Routes

The revised CSM for the Military Gas Station also refines the identification of potentially exposed receptor populations, receptor exposure points, and exposure routes for realistic scenarios based on specific site conditions. These components better reflect the likelihood and extent of human or ecological receptor contact with site-related contaminants. As described in Section 2, the Military Gas Station is entirely within the boundaries of the Base. Therefore, potential receptor groups are limited to onsite intrusive workers. There are no completed pathways to offsite receptors. Furthermore, the asphalt cover over much of the site, the depth to groundwater (35 feet bgs), and the lack of contamination in surface soils, prevents contact with contaminated soil or groundwater by current Base personnel.

The industrial nature of the site, and the pavement covering much of the site, precludes the existence of suitable wildlife habitat. No resident ecological receptors were identified for which soils and/or groundwater are likely contaminant exposure media. No exposure pathways involving potential offsite ecological receptors are or will be complete based on the lack of offsite contaminant migration.



Using the most conservative exposure assumptions appropriate for the Military Gas Station, the only realistic receptor that is likely to become exposed to site-related contaminants is the onsite intrusive worker involved in demolition, removal, and/or construction activities. Inhalation of VOCs (partitioning from either contaminated soil or groundwater) in ambient air at the site could result in a completed pathway for the onsite intrusive worker. However, as described in Section 4, there were no exceedances of soil TCLs, and soil gas concentrations are not expected to pose a significant inhalation risk to aboveground or intrusive workers. In addition, the water table at the site is substantially below the maximum construction depth, eliminating the risks posed by dermal contact or incidental ingestion of groundwater during construction activities. Therefore, these exposure pathways are either incomplete or are insignificant.

7.2.3 Summary of Exposure Pathway completion

Given the current and planned future uses of the Military Gas Station and the outcome of the chemical fate assessment presented in Section 6, only onsite intrusive workers could be exposed to site-related contamination in soils and soil gas during excavation activities. However, contaminant concentrations in these media are below Tier 1 action levels (Section 4), and do not pose a risk to potential receptors. Therefore, there are no currently completed exposure pathways at the Military Gas Station, and site data indicate that no exposure pathways will be completed in the future. As stated in Section 1, without a completed pathway through which a receptor will contact the contaminant(s), there is no risk posed by remaining levels of fuel hydrocarbon contamination at the site.

SECTION 8

SUMMARY AND CONCLUSIONS

Comparison of the COPC, electron acceptor, and biodegradation byproduct data for the Military Gas Station provides strong qualitative evidence of biodegradation of dissolved COPCs. Geochemical data strongly indicate that biodegradation of fuel hydrocarbons is occurring at the site, primarily via the processes of sulfate reduction and methanogenesis. The groundwater system appears to have more than enough capacity to facilitate biodegradation of all available contaminant mass dissolved in the groundwater and adsorbed to soil particles in the saturated zone. The measured decreases in dissolved contaminant concentrations in the source area prior to initiation of engineered remediation in 1994 or 1995 supports the effectiveness of natural attenuation processes in site groundwater. Direct evidence of the beneficial effects of biodegradation, in combination with the effects of air sparging and SVE, also is provided by the diminishment of the dissolved plume over time. As the contaminant source (residual LNAPL adsorbed to soil particles) is reduced over time due to biodegradation and SVE, dissolved contaminant concentrations in the source area also are reduced. The downgradient migration of the dissolved contaminants is severely restricted due to natural biodegradation. Available data indicate that the plume is entirely contained within the existing monitoring well network, and should not impact potential downgradient receptors. In addition, current and potential future onsite receptor exposure pathways are incomplete; therefore, the existing contamination does not pose a risk to potential receptors.

Because the dissolved plume is receding and is not currently impacting receptors, and because the contamination source will continue to decrease over time after the remediation system is shut off, RNA with long-term monitoring (LTM) and institutional controls is recommended for the Military Gas Station. A LTM plan is described in Section 9. The appropriateness of RNA with LTM is supported by the fact that only xylenes in source area well EA-4 exceeded the FDEP (1997) natural attenuation source default values (Table 4.2). Continued operation of the air sparging/SVE system is not necessary to minimize risks posed by site contamination, and shutoff of this system is recommended.

SECTION 9

LONG-TERM MONITORING PLAN

9.1 OVERVIEW

At the Military Gas Station, long-term monitoring (LTM) combined with RNA and institutional controls is recommended. The objectives of the LTM are as follows:

- To assess site conditions over time;
- To confirm the effectiveness of naturally occurring processes at reducing contaminant mass and minimizing contaminant migration;
- To evaluate the need for additional remediation.

The LTM plan consists of identifying groundwater sampling locations and developing a sampling and analysis strategy. The strategy described in this section is designed to assess the effectiveness of RNA through measurement of the reduction of contaminant mass, and the rate of groundwater remediation. In the event that data collected under this LTM program indicate that RNA is insufficient to be protective of human health and the environment, additional engineered controls to augment the beneficial effects of RNA may be necessary. A site-specific groundwater SAP and institutional control plan should be prepared prior to initiating the LTM program.

9.2 LONG-TERM GROUNDWATER MONITORING WELLS

Four monitoring wells should be included in the LTM program. These wells include two wells without hydrocarbon concentrations (one upgradient [EA-8] and one crossgradient [EA-11]) and two wells with historical hydrocarbon concentrations (EA-4 and EA-10) (Figure 5.1).

9.3 SAMPLING DURATION AND FREQUENCY

Up to nine years of monitoring may be required to accomplish the LTM objectives listed in Section 9.1. As described in Section 6.3.2, this is the estimated time frame for maximum dissolved xylenes concentrations to decrease below the Tier 1 TCL for this compound. Each of the LTM wells would be sampled annually for the first three years and biannually (every other year) for the following six years. The purpose of the long term sampling is to evaluate groundwater quality and to determine if the decreases observed in the dissolved COPC concentrations (Table 6.2) continue after the interim remediation system is shut down. If the data collected during this time period support the effectiveness of the selected remedial alternative at this site, it may be possible to

reduce or eliminate sampling. If the data collected at any time during the monitoring period indicate the need for additional remedial activities at the site, sampling frequency should be adjusted accordingly.

As new analytical results are obtained, they can be added to Figure 6.1 and the best-fit curves can be revised to refine decay rate estimates. The new decay rates can then be used to re-estimate the required duration of LTM, using the first-order rate equation contained in Appendix E.

9.4 ANALYTICAL PROTOCOL

All LTM wells will be sampled and analyzed to verify the effectiveness of naturally-occurring remediation processes at the site. At the beginning of each sampling event, water levels should be measured in all site monitoring wells. Groundwater samples collected from the LTM wells should be analyzed for the parameters listed in Table 9.1.

TABLE 9.1 ANALYTICAL PROTOCOL FOR GROUNDWATER ANALYSIS DURING LONG-TERM MONITORING

Military Gas Station Eglin AFB, Florida

ANALYTE	RECOMMENDED METHOD	WHERE ANALYZED
Ferrous Iron (Fe ⁺²)	Colorimetric, Hach Method 8146	Field
Sulfate (SO ₄ -2)	Colorimetric, Hach Method 8051	Field
Conductivity	Direct reading meter	Field
Oxygen	Direct reading meter	Field
рН	Direct reading meter	Field
Redox Potential	Direct reading meter	Field
Temperature	Direct reading meter	Field
BTEX ^{a/}	SW8021B	Fixed base lab
Naphthalene	SW8310	Fixed base lab
Total Lead	SW7421	Fixed base lab

Notes:

a/ BTEX = benzene, toluene, ethylbenzene, and total xylenes

SECTION 10

REFERENCES

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APPENDIX A LABORATORY ANALYTICAL DATA

VbENDIX V

LABORATORY ANALYTICAL DATA



AFCEE^Aromatic VOAs by Method 8020A with MTBE & TMBs Method 8020A

Client Name: Parsons Engineering Science

Client ID: MIL-EA-4 (0.00, 0.00)

LAB ID: 059721-0009-SA GRND-H20 Matrix:

Sampled: 31 MAR 98 Prepared: 06 APR 98 Received: 02 APR 98 Analyzed: 06 APR 98 02 APR 98 Authorized:

GCPID-H Dilution: 5.0 Instrument:

Parameter	Result	Qualifier	RL	MDL	Units
Benzene Toluene Ethylbenzene Xylenes (total)	ND ND 79 400	d d	10 10 10 10	0.28 0.75 0.27 0.75	ug/L ug/L ug/L ug/L

Surrogate	Recovery	Acceptable Ran		
a.a.a-Trifluorotoluene	98	*	44 - 165	
Fluorobenzene	86		44 - 165	

d = See Preferred Result on Other Column ND = Not Detected

Reported By: Shawn Hadley Approved By: Barbara Sullivan



AFCEE^Aromatic VOAs by Method 8020A (Second Column) Method 8020A

Client Name: Client ID: LAB ID:	Parsons Engineerin MIL-EA-4 059721-0009-SA	ng Scienc	ce (0.00,0	.00)			
Matrix: Authorized: Instrument:	GRND-H20 02 APR 98 GCPID-H	Prep	npled: 31 MAF bared: 06 APF ntion: 5.0	R 98 R 98	Received: Analyzed:		
Parameter		Result	Qualifier	RL	MDL	Units	
Benzene Toluene Ethylbenzene Xylenes (tota	1)	ND ND 76 400	M M	10 10 10 10	0.28 0.75 0.27 0.75	ug/L ug/L ug/L ug/L	
Surrogate			Recovery		Acceptable R	ange	
a.a.a-Trifluo Fluorobenzene	rotoluene		103 87	*	44 - 16 44 - 16		

M = Preferred Result
ND = Not Detected

Reported By: Shawn Hadley



AFCEE^Aromatic VOAs by Method 8020A with MTBE & TMBs Method 8020A

Client Name: Client ID: Parsons Engineering Science

MIL-EA-40 (0.00, 0.00)

LAB ID: 059721-0010-SA

Sampled: 31 MAR 98 Prepared: 06 APR 98 Dilution: 5.0 Matrix: GRND-H20 Received: 02 APR 98 Authorized: 02 APR 98 Analyzed: 06 APR 98 Instrument: GCPID-H

Parameter	Result	Qualifier	RL	MDL	Units
Benzene Toluene Ethylbenzene Xylenes (total)	ND ND 76 380	d M	10 10 10 10	0.28 0.75 0.27 0.75	ug/L ug/L ug/L ug/L

Surrogate	Recovery		Acceptable Range	
a.a.a-Trifluorotoluene	97	*	44 - 165	
Fluorobenzene	85		44 - 165	

d = See Preferred Result on Other Column

M = Preferred Result ND = Not Detected

Reported By: Shawn Hadley



AFCEE^Aromatic VOAs by Method 8020A (Second Column) Method 8020A

Client Name: Client ID: LAB ID:	Parsons Engineeri MIL-EA-40 059721-0010-SA	ng Sciend	ce (0.00,0.	00)		
Matrix: Authorized: Instrument:	GRND-H20 02 APR 98 GCPID-H	Prep	npled: 31 MAR bared: 06 APR ution: 5.0	98 98	Received: Analyzed:	02 APR 98 06 APR 98
Parameter		Result	Qualifier	RL	MDL	Units
Benzene Toluene Ethylbenzene Xylenes (tota	1)	ND ND 74 420	M d	10 10 10 10	0.28 0.75 0.27 0.75	ug/L ug/L ug/L ug/L
Surrogate			Recovery		Acceptable Ra	inge
a.a.a-Trifluon Fluorobenzene	rotoluene		102 85	z z	44 - 165 44 - 165	

d = See Preferred Result on Other Column
M = Preferred Result
ND = Not Detected

Reported By: Shawn Hadley



AFCEE^Aromatic VOAs by Method 8020A with MTBE & TMBs Method 8020A

Parsons Engineering Science

Client Name: Client ID: LAB ID: Mil-EA-5 059638-0003-SA (0.00, 0.00)

Sampled: 27 MAR 98 Prepared: 31 MAR 98 Dilution: 1.0 Received: 28 MAR 98 Analyzed: 31 MAR 98 GRND-H20 Matrix: 28 MAR 98

Authorized: GCPID-H Instrument:

Parameter	Result Qualifier	RL	MDL Units
Benzene Toluene Ethylbenzene Xylenes (total)	ND ND 1.4 JM 6.7 M	2.0 2.0 2.0 2.0	0.056 ug/L 0.15 ug/L 0.054 ug/L 0.15 ug/L
Surrogate	Recovery		Acceptable Range

Surrogate Acceptable Range a.a.a-Trifluorotoluene Fluorobenzene 44 - 165 44 - 165 89

 $\mbox{\bf J}$ = Result is detected below the reporting limit or is an estimated concentration. $\mbox{\bf M}$ = Preferred Result

ND = Not Detected

Reported By: Shawn Hadley



AFCEE^Aromatic VOAs by Method 8020A (Second Column) Method 8020A

Client Name: Client ID: LAB ID: Matrix: Authorized: Instrument:	Parsons Engineerin Mil-EA-5 059638-0003-SA GRND-H20 28 MAR 98 GCPID-H	Sam Prep	e (0.00,0.00 pled: 27 MAR 9 ared: 31 MAR 9 tion: 1.0	8	Received: 28 MAR 98 Analyzed: 31 MAR 98
Parameter		Result	Qualifier	RL	MDL Units
Benzene Toluene Ethylbenzene Xylenes (tota		ND ND 1.5 6.9	Jd d	2.0 2.0 2.0 2.0	0.056 ug/L 0.15 ug/L 0.054 ug/L 0.15 ug/L
Surrogate			Recovery		Acceptable Range
a.a.a-Trifluo Fluorobenzene			98 91	X	44 - 165 44 - 165

Reported By: Shawn Hadley

d = See Preferred Result on Other Column J = Result is detected below the reporting limit or is an estimated concentration. ND = Not Detected $\,$



AFCEE^Aromatic VOAs by Method 8020A with MTBE & TMBs Method 8020A

Client	Name:	Parsons	Engineering	Science
• •				

Client ID: Mil-EA-8

(0.00, 0.00)

LAB ID: 059638-0004-SA

Sampled: 27 MAR 98 Prepared: 31 MAR 98 Dilution: 1.0 Received: 28 MAR 98 Analyzed: 31 MAR 98 GRND-H20 28 MAR 98 Matrix: Authorized:

GCPID-H Instrument:

Parameter	Result Qualifier	RL	MDL Units
Benzene Toluene Ethylbenzene Xylenes (total)	ND ND 0.099 JM 4.7 d	2.0 2.0 2.0 2.0	0.056 ug/L 0.15 ug/L 0.054 ug/L 0.15 ug/L
Surrogate	Recovery		Acceptable Range

a,a,a-Trifluorotoluene Fluorobenzene 44 - 165 44 - 165 92

d = See Preferred Result on Other Column
J = Result is detected below the reporting limit or is an estimated concentration.
M = Preferred Result

ND = Not Detected

Reported By: Shawn Hadley



AFCEE^Aromatic VOAs by Method 8020A (Second Column) Method 8020A

Client Name: Client ID: LAB ID:	Parsons Engineeri Mil-EA-8 059638-0004-SA	ng Sciend	ce (0.00,0.0	0)	
Matrix: Authorized: Instrument:	GRND-H20 28 MAR 98 GCPID-H	Prep	npled: 27 MAR 9 Dared: 31 MAR 9 Darion: 1.0	98 98	Received: 28 MAR 98 Analyzed: 31 MAR 98
Parameter		Result	Qualifier	RL	MDL Units
Benzene Toluene Ethylbenzene Xylenes (tota	1)	ND ND 1.1 4.7	Jd M	2.0 2.0 2.0 2.0	0.056 ug/L 0.15 ug/L 0.054 ug/L 0.15 ug/L
Surrogate			Recovery		Acceptable Range
a.a.a-Trifluo Fluorobenzene	rotoluene		94 89	*	44 - 165 44 - 165

Reported By: Shawn Hadley

d = See Preferred Result on Other Column J = Result is detected below the reporting limit or is an estimated concentration. M = Preferred Result ND = Not Detected



AFCEE^Aromatic VOAs by Method 8020A with MTBE & TMBs Method 8020A

074	Mama.	Danasas	Engineening	Coionco
Cilent	Name:	Parsons	Engineering	or i clice

(0.00, 0.00)Client ID:

LAB ID:

Mil-EA-10 059638-0002-SA GRND-H20 28 MAR 98 Sampled: 27 MAR 98 Prepared: 31 MAR 98 Dilution: 1.0 Received: 28 MAR 98 Analyzed: 31 MAR 98 Matrix: Authorized:

GCPID-H Instrument:

Parameter	Result Qualifier	RL	MDL Units
Benzene Toluene Ethylbenzene Xylenes (total)	ND ND 2.3 M 11 M	2.0 2.0 2.0 2.0	0.056 ug/L 0.15 ug/L 0.054 ug/L 0.15 ug/L

Surrogate	Recovery		Acceptable Range
a,a,a-Trifluorotoluene	100	×	44 - 165
Fluorobenzene	92		44 - 165

M = Preferred Result ND = Not Detected

Reported By: Shawn Hadley

Approved By: Audrey Cornell



AFCEE^Aromatic VOAs by Method 8020A (Second Column) Method 8020A

Client Name:	Parsons Engineering Science	
Client ID:	Mil-EA-10	(0.00, 0.00)
LAD TD.	050630 0003 CA	

LAB ID: Matrix:	059638-0002-5A GRND-H2O	Sampled: 27 MAR 98	Received: 28 MAR 98
Authorized:	28 MAR 98	Prepared: 31 MAR 98	Analyzed: 31 MAR 98

Instrument:	GCPID-H	Dilution:	1.0

Parameter	Result (Qualifier	RL	MDL	Units
Benzene Toluene Ethylbenzene Xylenes (total)	ND ND 2.3 11	d d	2.0 2.0 2.0 2.0	0.056 0.15 0.054 0.15	ug/L ug/L ug/L ug/L

Surrogate	Recovery		Acceptable Range
a,a,a-Trifluorotoluene	100	X	44 - 165
Fluorobenzene	92	X	44 - 165

 $d = See \ Preferred \ Result on \ Other \ Column \ ND = Not \ Detected$

Reported By: Shawn Hadley



AFCEE Polynuclear Aromatic Hydrocarbons, HPLC Method 8310

P7	Mama	Dancene	Engineening	Caianaa
Litent	name:	Parsons	Engineering	oc rence

MIL-EA-4 059721-0009-SA (0.00, 0.00)Client ID:

LAB ID:

Sampled: 31 MAR 98 Prepared: 07 APR 98 Dilution: 0.98 Received: 02 APR 98 Analyzed: 22 APR 98 GRND-H20 Matrix: 02 APR 98 Authorized:

HPLC-Y Instrument:

211001 41101101				
Parameter	Result Qualifier	RL	MDL	Units
Acenaphthene Acenaphthylene Anthracene Benzo(a)anthracene Benzo(b)fluoranthene Benzo(g,h,i)perylene Benzo(k)fluoranthene Chrysene Dibenz(a,h)anthracene Fluorene Fluoranthene Indeno(1,2,3-cd)pyrene Naphthalene Phenanthrene Pyrene	ND N	0.98 0.98 0.098 0.13 0.23 0.18 0.20 0.17 0.20 0.29 0.20 0.42 0.98 0.20	0.091 0.039 0.029 0.059 0.064 0.058 0.070 0.058 0.072 0.074 0.034 0.035 0.054 0.058	ug/L ug/L ug/L ug/L ug/L ug/L ug/L ug/L
Surrogate	Recovery	Ac	ceptable Ra	nge

Terphenyl-d14 106 X 25 - 157

 $\begin{array}{ll} \textbf{d} &= \text{See Preferred Result on Other Column} \\ \textbf{J} &= \text{Result is detected below the reporting limit or is an estimated concentration.} \end{array}$

ND = Not Detected

Reported By: Blake Besser



AFCEE Polynuclear Aromatic Hydrocarbons, HPLC (Second Column) Method 8310

(0.00, 0.00)

Client Name: Parsons Engineering Science

Client ID: MIL-EA-4

LAB ID: 059721-0009-SA

Matrix: GRND-H20 Sampled: 31 MAR 98 Received: 02 APR 98 Authorized: 02 APR 98 Prepared: 07 APR 98 Analyzed: 22 APR 98

Instrument: HPLC-Y Dilution: 0.98

Parameter	Result Qualifier	RL	MDL Units
Acenaphthene Acenaphthylene Anthracene Benzo(a)anthracene Benzo(b)fluoranthene Benzo(g,h,i)perylene Benzo(k)fluoranthene Chrysene Dibenz(a,h)anthracene Fluorene Fluoranthene Indeno(1,2,3-cd)pyrene Naphthalene Phenanthrene	Result Qualifier ND	0.98 0.98 0.098 0.13 0.23 0.18 0.20 0.17 0.20 0.29 0.20 0.42 0.98 0.20	MDL Units 0.091 ug/L 0.039 ug/L 0.029 ug/L 0.059 ug/L 0.058 ug/L 0.058 ug/L 0.070 ug/L 0.058 ug/L 0.072 ug/L 0.074 ug/L 0.034 ug/L 0.035 ug/L 0.054 ug/L 0.058 ug/L
Pyrene	ND	0.20	0.042 ug/L

Surrogate Recovery Acceptable Range
Terphenyl-d14 108 % 25 - 157

ND = Not Detected

Reported By: Blake Besser

 $^{{\}tt J}$ = Result is detected below the reporting limit or is an estimated concentration. ${\tt M}$ = Preferred Result



AFCEE Polynuclear Aromatic Hydrocarbons, HPLC Method 8310

Client	Namo ·	Darcone	Fngineering	Science

Client ID: (0.00, 0.00)

MIL-EA-40 059721-0010-SA GRND-H20 LAB ID: Matrix: Sampled: 31 MAR 98 Prepared: 07 APR 98 Dilution: 1.0 Received: 02 APR 98 Analyzed: 22 APR 98 Authorized: 02 APR 98

HPLC-Y Instrument:

Parameter	Result Qualifier	RL	MDL Units
Acenaphthene Acenaphthylene Anthracene Benzo(a)anthracene Benzo(b)fluoranthene Benzo(b)fluoranthene Benzo(k)fluoranthene Chrysene Dibenz(a,h)anthracene Fluorene Fluoranthene Indeno(1,2,3-cd)pyrene Naphthalene Phenanthrene Pyrene	ND N	1.0 1.0 0.10 0.13 0.23 0.18 0.20 0.17 0.20 0.31 0.20 0.20 0.44 1.0 0.20	0.095 ug/L 0.041 ug/L 0.031 ug/L 0.061 ug/L 0.066 ug/L 0.060 ug/L 0.072 ug/L 0.074 ug/L 0.078 ug/L 0.036 ug/L 0.037 ug/L 0.036 ug/L 0.037 ug/L 0.056 ug/L 0.044 ug/L

Surrogate Recovery Acceptable Range Terphenyl-d14 98 X 25 - 157

 $d = See \ Preferred \ Result on \ Other \ Column \ J = Result is detected below the reporting limit or is an estimated concentration.$ $ND = Not \ Detected$

Reported By: Blake Besser



AFCEE Polynuclear Aromatic Hydrocarbons, HPLC (Second Column) Method 8310

Client	Name:	Parsons	Engineering	Science
CITCHE	Hume.	1 01 30113	Ling incci ing	JC I CIICE

Parameter

Acenaphthene

Terphenyl-d14

Acenaphthylene Anthracene

Client ID: MIL-EA-40 (0.00, 0.00)

LAB ID: 059721-0010-SA Sampled: 31 MAR 98 Prepared: 07 APR 98 Matrix: GRND-H20 Received: 02 APR 98 02 APR 98 Authorized: Analyzed: 22 APR 98 HPLC-Y Dilution: 1.0 Instrument:

Result Qualifier

ND

ND

RL

1.0

1.0

X

MDL

0.095

0.041

25 - 157

Units

ug/L

ug/L

Anthracene	ND	0.10	0.031	ug/L
Benzo(a)anthracene	ND	0.13	0.061	ug/L
Benzo(a)pyrene	ND	0.23	0.066	ug/L
Benzo(b)fluoranthene	ND	0.18	0.060	ug/L
Benzo(g.h.i)perylene	ND	0.20	0.072	ug/L
Benzo(k)fluoranthene	ND	0.17	0.060	ug/L
Chrysene	ND	0.20	0.074	ug/L
Dibenz(a,h)anthracene	ND	0.31	0.078	ug/L
Fluorene	0.085 JM	0.20	0.036	ug/L
Fluoranthene	ND	0.20	0.037	ug/L
Indeno(1,2,3-cd)pyrene	ND	0.44	0.056	ug/L
Naphthalene	27 M	1.0	0.060	ug/L
Phenanthrene	ND	0.20	0.031	ug/L
Pyrene	ND	0.20	0.044	ug/L
Surrogate	Recovery Acceptable Range			inge

98

ND = Not Detected

Reported By: Blake Besser

J = Result is detected below the reporting limit or is an estimated concentration. M = Preferred Result



AFCEE Polynuclear Aromatic Hydrocarbons, HPLC Method 8310

C7	Mama.	Dancone	Engineering	Scriance
LITERE	Name:	Parsons	end neer mu	2C LELICE

Client ID: Mil-EA-10 (0.00, 0.00)

059638-0002-SA

LAB ID: Matrix: Sampled: 27 MAR 98 Prepared: 01 APR 98 Dilution: 1.0 Received: 28 MAR 98 Analyzed: 18 APR 98 GRND-H20 Authorized: 28 MAR 98

HPLC-Q Instrument:

Parameter	Result Qualifier	RL	MDL	Units	
Acenaphthene Acenaphthylene Anthracene Benzo(a)anthracene Benzo(b)fluoranthene Benzo(g,h,i)perylene Benzo(k)fluoranthene Chrysene Dibenz(a,h)anthracene Fluorene Fluoranthene Indeno(1,2,3-cd)pyrene Naphthalene Phenanthrene Pyrene	ND N	1.0 1.0 0.10 0.13 0.23 0.18 0.20 0.17 0.20 0.30 0.20 0.44 1.0 0.20	0.094 0.041 0.030 0.061 0.066 0.072 0.060 0.074 0.077 0.036 0.037 0.056 0.060	ug/L ug/L ug/L ug/L ug/L ug/L ug/L ug/L	
Cumpagata	Pacayany	٨٥	contable Day	ngo	

Surrogate Recovery Acceptable Range 99 X 25 - 157 Terphenyl-d14

d = See Preferred Result on Other Column J = Result is detected below the reporting limit or is an estimated concentration.

ND = Not Detected

Reported By: Blake Besser



AFCEE Polynuclear Aromatic Hydrocarbons, HPLC (Second Column) Method 8310

Client Name: Parsons Engineering Science Client ID: Mil-EA-10 (0.00, 0.00)

LAB ID:

059638-0002-SA GRND-H20 Sampled: 27 MAR 98 Prepared: 01 APR 98 Dilution: 1.0 Received: 28 MAR 98 Analyzed: 18 APR 98 Matrix: Authorized: 28 MAR 98

Instrument: HPLC-Q

Parameter	Result Qualifier	RL	MDL Unit	S
Acenaphthene Acenaphthylene Anthracene Benzo(a)anthracene Benzo(b)fluoranthene Benzo(g,h.i)perylene Benzo(k)fluoranthene Chrysene Dibenz(a,h)anthracene Fluorene Fluoranthene Indeno(1,2,3-cd)pyrene Naphthalene Phenanthrene Pyrene	ND N	1.0 1.0 0.10 0.13 0.23 0.18 0.20 0.17 0.20 0.30 0.20 0.20 0.44 1.0 0.20 0.20	0.094 ug/ 0.041 ug/ 0.030 ug/ 0.061 ug/ 0.066 ug/ 0.072 ug/ 0.074 ug/ 0.077 ug/ 0.036 ug/ 0.037 ug/ 0.056 ug/ 0.056 ug/ 0.056 ug/ 0.056 ug/	
Surrogate	Recovery	A	cceptable Range	
Tambanyl dlA	102	٠	25 - 157	

^{25 - 157} Terphenyl-d14 102

 ${\tt J}$ = Result is detected below the reporting limit or is an estimated concentration. ${\tt M}$ = Preferred Result ${\tt ND}$ = Not Detected

Reported By: Blake Besser



(0.00, 0.00)

Client Name: Parsons Engineering Science

Client ID: MIL-EA-4

LAB ID: 059721-0009-SA

Sampled: 31 MAR 98 Prepared: 13 APR 98 Dilution: 1.0 GRND-H20 02 APR 98 Matrix: Received: 02 APR 98 Analyzed: 14 APR 98 Authorized:

Instrument: GCFID-I

Parameter Result Qualifier RL MDL Units

EDB (1.2-Dibromoethane) ND 0.020 0.0060 ug/L

Surrogate Recovery Acceptable Range

1.1.1.2-Tetrachloroethane 114 X 80 - 120

ND = Not Detected



Client Name:

Parsons Engineering Science

Client ID:

MIL-EA-40

(0.00, 0.00)

LAB ID: Matrix:

059721-0010-SA

GRND-H20

Received: 02 APR 98

Authorized: Instrument: 02 APR 98 GCFID-I

Sampled: 31 MAR 98 Prepared: 13 APR 98 Dilution: 1.0

Analyzed: 14 APR 98

Parameter

Result Qualifier

RL MDL

Units

EDB (1.2-Dibromoethane)

ND

0.020

X

0.0060 ug/L

Surrogate

Recovery

Acceptable Range

1.1.1.2-Tetrachloroethane

106

80 - 120

ND = Not Detected

Reported By: Quanterra-Tampa



Client Name: Client ID: Parsons Engineering Science

Mil-EA-5 (0.00, 0.00)

059638-0003-SA LAB ID:

Sampled: 27 MAR 98 Prepared: 03 APR 98 Dilution: 1.0 Received: 28 MAR 98 GRND-H20 Matrix: Analyzed: 04 APR 98 28 MAR 98 Authorized:

GCFID-I Instrument:

MDL RL Units Parameter Result Qualifier

0.020 0.0060 ug/L ND EDB (1.2-Dibromoethane)

Acceptable Range Recovery Surrogate

X 80 - 120 104 1.1.1.2-Tetrachloroethane

ND = Not Detected



Client Name: Parsons Engineering Science Client ID: Mil-EA-8 (0.00, 0.00)

LAB ID:

059638-0004-SA GRND-H20 Sampled: 27 MAR 98 Prepared: 03 APR 98 Dilution: 1.0 Received: 28 MAR 98 Analyzed: 04 APR 98 Matrix: 28 MAR 98 Authorized:

GCFID-I Instrument:

RL Result Qualifier MDL Units Parameter

ND 0.020 0.0060 ug/L EDB (1.2-Dibromoethane)

Acceptable Range Surrogate Recovery

80 - 120 × 1,1,1,2-Tetrachloroethane 111

ND = Not Detected



(0.00, 0.00)

Client Name: Client ID: LAB ID: Parsons Engineering Science

Mil-EA-10

059638-0002-SA

Matrix:

GRND-H20

28 MAR 98 Authorized:

Sampled: 27 MAR 98 Prepared: 03 APR 98 Received: 28 MAR 98 Analyzed: 04 APR 98

Dilution: 1.0 GCFID-I Instrument:

MDL Result Qualifier RL Units Parameter

0.020 0.0060 ug/L EDB (1,2-Dibromoethane) ND

Acceptable Range Recovery Surrogate

80 - 120 104 × 1.1.1.2-Tetrachloroethane

ND = Not Detected



Method FL-PRO - TPH (C8-C40) Method FL-PRO

(0.00, 0.00)

Client Name: Parsons Engineering Science

Client ID: MIL-EA-4

LAB ID: 059721-0009-SA

Sampled: 31 MAR 98 Prepared: 06 APR 98 Dilution: 1.0 GRND-H20 Matrix: Received: 02 APR 98 Authorized: 02 APR 98 Analyzed: 08 APR 98

Instrument: GCFID-I

Parameter Result Qualifier RL MDL Units TPH (C8-C40) 3.1 0.50 0.19 mg/L Surrogate Recovery Acceptable Range o-Terphenyl Nonatriacontane * 33 - 162 10 - 109 103 81

Reported By: Quanterra-Tampa



Method FL-PRO - TPH (C8-C40) Method FL-PRO

(0.00, 0.00)

Client Name: Parsons Engineering Science
Client ID: MIL-EA-40
LAB ID: 059721-0010-SA
Matrix: GRND-H20 Sampl
Authorized: 02 APR 98 Prepar Sampled: 31 MAR 98 Prepared: 06 APR 98 Dilution: 1.0 Received: 02 APR 98 Analyzed: 08 APR 98

GCFID-I Instrument:

Parameter Result Qualifier RL MDL Units

TPH (C8-C40) 4.4 0.50 0.19 mg/L

Surrogate Recovery Acceptable Range

o-Terphenyl Nonatriacontane 33 - 162 10 - 109 108 · * 98

Reported By: Quanterra-Tampa



Method FL-PRO - TPH (C8-C40) Method FL-PRO

(0.00, 0.00)

Client Name: Client ID: Parsons Engineering Science

Mil-EA-10

LAB ID: 059638-0002-SA

Sampled: 27 MAR 98 Prepared: 02 APR 98 Dilution: 1.0 Received: 28 MAR 98 Analyzed: 03 APR 98 GRND-H20 Matrix: 28 MAR 98 Authorized:

GCFID-I Instrument:

MDL Units Result Qualifier RL Parameter 0.50 0.19 mg/L 0.81 TPH (C8-C40)

Recovery Acceptable Range Surrogate

33 - 162 10 - 109 102 o-Terphenyl Nonatriacontane 39



Client Name: Client ID: LAB ID:

Parsons Engineering Science

(0.00, 0.00)

MIL-EA-4 059721-0009-SA

Matrix:

GRND-H20

Sampled: 31 MAR 98 Prepared: N/A

Authorized: Instrument: 02 APR 98 GCFID-K1A

Dilution: 50

Received: 02 APR 98 Analyzed: 13 APR 98

Parameter

Result Qualifier

RL

MDL Units

Methane

1700

В

25

2.6

ug/L

B = Compound is also detected in the blank.

Reported By: Quanterra-Austin



Parsons Engineering Science Client Name:

Client ID: MIL-EA-40 (0.00, 0.00)

059721-0010-SA LAB ID:

Sampled: 31 MAR 98 Prepared: N/A Dilution: 50 GRND-H20 Matrix: Received: 02 APR 98 02 APR 98 Authorized: Analyzed: 13 APR 98

Instrument: GCFID-K1A

Parameter Result Qualifier MDL RL Units

Methane 1500 В 25 2.6 ug/L

B = Compound is also detected in the blank.

Reported By: Quanterra-Austin

Client Name: Parsons Engineering Science Client ID: Mil-EA-5

(0.00.0.00)

LAB ID: Matrix: 059638-0003-SA GRND-H20

Received: 28 MAR 98

Authorized:

Parameter

28 MAR 98

Sampled: 27 MAR 98 Prepared: N/A

Analyzed: 07 APR 98

Instrument: GCFID-K1A Dilution: 1.0

Result Qualifier

MDL

Units

Methane

0.54 B

0.50

RL

0.052

ug/L

= Compound is also detected in the blank.

Reported By: Quanterra-Austin

Client Name: Client ID: LAB ID:

Parsons Engineering Science

Mil-EA-8

(0.00.0.00)

059638-0004-SA

Matrix:

GRND-H20 28 MAR 98 Sampled: 27 MAR 98 Prepared: N/A Dilution: 1.0

JB

Authorized:

Parameter

GCFID-K1A

Received: 28 MAR 98 Analyzed: 07 APR 98

Instrument:

Result Qualifier

MDL Units

Methane

0.38

0.50

RL

0.052

ug/L

B = Compound is also detected in the blank.
 J = Result is detected below the reporting limit or is an estimated concentration.

Parsons Engineering Science Client Name:

GRND-H20

Client ID: Mil-EA-10

059638-0002-SA LAB ID:

Matrix:

(0.00, 0.00)

Authorized:

Sampled: 27 MAR 98 Prepared: N/A Dilution: 1.0 Received: 28 MAR 98 Analyzed: 07 APR 98 28 MAR 98 GCFID-KlA Instrument:

Result Oualifier RL MDL Units Parameter

0.50 0.052 Methane 1.2 В ug/L

- Compound is also detected in the blank.

Reported By: Quanterra-Austin



AFCEE Total Metals

Client Name:

Parsons Engineering Science

Client ID: LAB ID:

Mi1-EA-04

059619-0006-SA

Matrix:

GRND-H20

Authorized:

27 MAR 98

(0.00, 0.00)

Sampled: 26 MAR 98 Prepared: See Below

Received: 27 MAR 98 Analyzed: See Below

Parameter

Result

Qual Dil

1.0

MDL

Rep Lim Units

Method

Prepared Analyzed Date Date

Lead

0.015

0.0010

0.0050mg/L

SW7421

31 MAR 98 01 APR 98

Reported By: Robin Proctor



AFCEE Total Metals

Client Name: Client ID: LAB ID: Matrix:

Authorized:

Parsons Engineering Science Mil-EA-40

059619-0007-SA

GRND-H20

27 MAR 98

(0.00, 0.00)

Sampled: 26 MAR 98 Prepared: See Below

Received: 27 MAR 98 Analyzed: See Below

Parameter

Result Qual Dil MDL

Rep Lim Units

Method

Prepared Analyzed Date Date

Lead

0.015

1.0

0.0010

0.0050mg/L

SW7421

31 MAR 98 01 APR 98

Reported By: Robin Proctor



AFCEE Total Metals

Client Name:

Parsons Engineering Science

Client ID: LAB ID:

(0.00, 0.00)

Matrix:

Mil-EA-10 059638-0002-SA GRND-H20 28 MAR 98

Authorized:

Sampled: 27 MAR 98 Prepared: See Below

Received: 28 MAR 98 Analyzed: See Below

Parameter

Qual Dil Result

MDL

Rep Lim Units Method Date

Prepared Analyzed Date

Lead

0.011

0.0010 1.0

0.0050mg/L

SW7421

31 MAR 98 01 APR 98

Reported By: Robin Proctor

Approved By: Kathy Wakeman



AFCEE Dissolved Metals

Client Name:

Parsons Engineering Science

Client ID:

Mil-EA-04

LAB ID:

059619-0006-SA

(0.00, 0.00)

Matrix: Authorized: GRND-H20 27 MAR 98

Sampled: 26 MAR 98

Prepared: See Below

Received: 27 MAR 98

Analyzed: See Below

Parameter

Result

Qual Dil

MDL

Rep Lim Units

Method

Prepared Analyzed Date

Date

Lead

0.017

1.0 0.0010 0.0050mg/L

SW7421

03 APR 98 06 APR 98

Reported By: Robin Proctor



AFCEE Dissolved Metals

Client Name:

Parsons Engineering Science

Client ID:

Mil-EA-40

LAB ID:

059619-0007-SA

Matrix: Authorized: GRND-H20 27 MAR 98

(0.00, 0.00)

0.0010

Sampled: 26 MAR 98 Prepared: See Below

Received: 27 MAR 98 Analyzed: See Below

Parameter

Result Qual Dil MDL

Rep Lim Units

Method

Prepared Analyzed Date

Date

Lead

0.017

1.0

0.0050mg/L

SW7421

03 APR 98 06 APR 98

Reported By: Robin Proctor



AFCEE Dissolved Metals

Client Name:

Parsons Engineering Science

Client ID:

LAB ID:

Mil-EA-10 059638-0002-SA GRND-H20

(0.00, 0.00)

Matrix:

Sampled: 27 MAR 98 Prepared: See Below

Received: 28 MAR 98

Authorized:

28 MAR 98

Analyzed: See Below

Parameter

Result Qual Dil MDL

Rep Lim Units Method Prepared Analyzed Date

Date

Lead

0.019

1.0 0.0010 0.0050mg/L

SW7421

03 APR 98 06 APR 98

Reported By: Robin Proctor

Approved By: Kathy Wakeman



Client Name: Client ID: LAB ID:

Parsons Engineering Science

Mil-EA-04

059619-0006-SA

Matrix:

GRND-H20

Authorized:

27 MAR 98

Sampled: 26 MAR 98

(0.00, 0.00)

Prepared: See Below

Received: 27 MAR 98

Analyzed: See Below

Prepared Analyzed Rep Lim Units MDL Method Date Date Parameter Result Qual Dil

NA 0.040 0.50 mg/L 9056 27 MAR 98 4.3 1.0 Nitrate as N

Reported By: Patty Jungk



Client Name:

Parsons Engineering Science

Client ID:

Authorized:

Mil-EA-40

LAB ID: Matrix:

059619-0007-SA

GRND-H20

(0.00, 0.00)

27 MAR 98

Sampled: 26 MAR 98 Prepared: See Below

Received: 27 MAR 98 Analyzed: See Below

Parameter

Result

Qual Dil MDL Rep Lim Units Method

Date

Date

Nitrate as N

4.3

0.040 1.0

0.50 mg/L

9056

NA

27 MAR 98

Prepared Analyzed

Reported By: Patty Jungk



Client Name:

Parsons Engineering Science

Client ID:

Mil-EA-5

(0.00, 0.00)

LAB ID:

059638-0003-SA GRND-H20

Matrix:

Sampled: 27 MAR 98

Received: 28 MAR 98 Analyzed: See Below

Authorized:

28 MAR 98

Prepared: See Below

Parameter

Qual Dil Result

MDL

Rep Lim Units Method Prepared Analyzed Date Date

Nitrate as N

3.7

0.040 1.0

0.50 mg/L

9056

NA

29 MAR 98

Reported By: Patty Jungk



Client Name:

Parsons Engineering Science Mil-EA-8 059638-0004-SA GRND-H20 Samp 28 MAR 98 Prep

Client ID:

LAB ID:

Matrix:

1.4

Authorized:

(0.00, 0.00)

Sampled: 27 MAR 98 Prepared: See Below

Received: 28 MAR 98 Analyzed: See Below

Parameter

Nitrate as N

Result Qual Dil

1.0

MDL

0.040

Rep Lim Units 0.50 mg/L

9056

Method

NA

Prepared Analyzed Date Date

29 MAR 98

Reported By: Patty Jungk



Client Name: Client ID: LAB ID:

Parsons Engineering Science

Mil-EA-10

(0.00, 0.00)

059638-0002-SA

Matrix:

GRND-H20

Received: 28 MAR 98

Authorized:

28 MAR 98

Sampled: 27 MAR 98 Prepared: See Below

Analyzed: See Below

Parameter

Result Qual Dil MDL

Method Rep Lim Units

Prepared Analyzed Date Date

Nitrate as N

1.2

0.040 1.0

0.50 mg/L

9056

28 MAR 98 NA

Reported By: Patty Jungk



AFCEE^Aromatic Volatile Organics with MTBE Method 8020A

Parsons Engineering Science Client Name:

MIL-B-S25 059724-0008-SA Client ID: (0.00, 0.00)

LAB ID:

Sampled: 29 MAR 98 Prepared: 07 APR 98 Dilution: 1.0 Received: 02 APR 98 Matrix: SOIL 02 APR 98 Analyzed: 07 APR 98 Authorized:

GCPID-H Instrument:

Parameter	Result	Qualifier	RL	MDL	Units
Benzene	ND		0.0061	0.00061	mg/kg
Toluene	ND		0.0061	0.0015	mg/kg
Ethylbenzene	ND		0.0024	0.00061	mg/kg
Xylenes (total)	ND		0.0061	0.0018	mg/kg
Methyl-tert-butyl ether	ND		0.0061	0.0018	mg/kg

Surrogate	Recovery		Acceptable Range
a,a,a-Trifluorotoluene	102	X	34 - 175
Fluorobenzene	108	X	34 - 175

Percent moisture is 17.7%. All results and limits are reported on a dry weight basis.

ND = Not Detected

Reported By: Shawn Hadley Approved By: Barbara Sullivan



AFCEE^Aromatic Volatile Organics with MTBE Method 8020A

Parsons Engineering Science MIL-K-S15 059724-0009-SA Client Name: Client ID: (0.00, 0.00)

LAB ID:

Sampled: 30 MAR 98 Prepared: 07 APR 98 Dilution: 1.0 Received: 02 APR 98 Analyzed: 07 APR 98 SOIL Matrix: 02 APR 98 Authorized:

Instrument: GCPID-H

Parameter	Result	Qualifier	RL	MDL	Units
Benzene Toluene Ethylbenzene Xylenes (total) Methyl-tert-butyl ether	ND ND ND ND ND		0.0055 0.0055 0.0022 0.0055 0.0055	0.00055 0.0013 0.00055 0.0016 0.0016	mg/kg mg/kg mg/kg

urrogate Recovery			Acceptable Range		
a.a.a-Trifluorotoluene	100	X	34 - 175		
Fluorobenzene	100		34 - 175		

Percent moisture is 9.0%.

All results and limits are reported on a dry weight basis.

ND = Not Detected

Reported By: Shawn Hadley

Approved By: Barbara Sullivan



AFCEE^Aromatic Volatile Organics with MTBE Method 8020A

Client Name:

Parsons Engineering Science MIL-V-S26 Client ID: (0.00, 0.00)

LAB ID: 059724-0010-SA

Sampled: 29 MAR 98 Prepared: 07 APR 98 Dilution: 1.0 Matrix: SOIL Received: 02 APR 98 Authorized: 02 APR 98 Analyzed: 07 APR 98

GCPID-H . Instrument:

Parameter Result Qualifier RL MDL Units Benzene ND 0.0061 0.00060 mg/kg Toluene ND 0.0061 0.0015 mg/kg Ethylbenzene 0.0050 M 0.0024 0.00060 mg/kg Xylenes (total) 0.012 d 0.0061 0.0018 mg/kg Methyl-tert-butyl ether ND 0.0061 0.0018 mg/kg

Surrogate Recovery Acceptable Range a.a.a-Trifluorotoluene 34 - 175 34 - 175 101 Fluorobenzene 103 *

Percent moisture is 17.4%. All results and limits are reported on a dry weight basis.

d = See Preferred Result on Other Column

M = Preferred Result ND = Not Detected

Reported By: Shawn Hadley

Approved By: Barbara Sullivan



AFCEE^Aromatic Volatile Organics with MTBE (Second Column) Method 8020A

Client Name: Parsons Engineering Science

Client ID: MIL-V-S26 (0.00, 0.00)

059724-0010-SA LAB ID:

Sampled: 29 MAR 98 Prepared: 07 APR 98 Dilution: 1.0 Received: 02 APR 98 Matrix: SOIL Authorized: 02 APR 98 Analyzed: 07 APR 98

Instrument: GCPID-H

Parameter	Result Qualifier	RL	MDL Units
Benzene Toluene Ethylbenzene Xylenes (total) Methyl-tert-butyl ether	ND	0.0061	0.00060 mg/kg
	ND	0.0061	0.0015 mg/kg
	0.0050 d	0.0024	0.00060 mg/kg
	0.012 M	0.0061	0.0018 mg/kg
	ND	0.0061	0.0018 mg/kg

Surrogate	Recovery	Acce	ptable Range
a,a,a-Trifluorotoluene	101	· -	34 - 175
Fluorobenzene	104		34 - 175

Percent moisture is 17.4%. All results and limits are reported on a dry weight basis.

d = See Preferred Result on Other Column

M = Preferred Result ND = Not Detected

Reported By: Shawn Hadley

Approved By: Barbara Sullivan



AFCEE Polynuclear Aromatic Hydrocarbons, HPLC Method 8310

RL

X

MDL

22 - 167

Units

Client Name: Parsons Engineering Science Client ID: MIL-B-S25 LAB ID: 059724-0008-SA

(0.00, 0.00)

Parameter

Terphenyl-d14

Sampled: 29 MAR 98 Prepared: 06 APR 98 Dilution: 1.0 Received: 02 APR 98 Analyzed: 23 APR 98 Matrix: SOIL 02 APR 98 HPLC-Y Authorized:

Result Qualifier

Instrument:

	•			
Acenaphthene	ND	0.24	0.044 mg/k	g
Acenaphthylene	ND	0.24	0.060 mg/k	g
Anthracene	ND	0.024	0.0032 mg/k	g
Benzo(a)anthracene	ND	0.024	0.0041 mg/k	g
Benzo(a)pyrene	ND	0.018	0.0027 mg/k	g
Benzo(b)fluoranthene	ND	0.015	0.0030 mg/k	g
Benzo(g,h,i)perylene	ND	0.061	0.0036 mg/k	ğ
Benzo(k)fluoranthene	ND	0.013	0.0043 mg/k	g
Chrysene	ND	0.049	0.0024 mg/k	ğ
Dibenz(a,h)anthracene	ND	0.024	0.0041 mg/k	ā
Fluorene	ND	0.049	0.0078 mg/k	ă
Fluoranthene	ND	0.049	0.0029 mg/k	a
Indeno(1,2,3-cd)pyrene	ND	0.036	0.0036 mg/k	a
Naphthalene	ND	0.24	0.056 mg/k	э П
Phenanthrene	ND	0.049	0.0067 mg/k	σ.
		0.049	0.0032 mg/k	9
Pyrene	ND	0.049	0.0032 mg/K	9
Surrogate	Recovery	Acce	ptable Range	

100

Percent moisture is 17.7%. All results and limits are reported on a dry weight basis.

ND = Not Detected

Reported By: Blake Besser Approved By: Audrey Cornell



AFCEE Polynuclear Aromatic Hydrocarbons, HPLC Method 8310

Client Name: Parsons Engineering Science

(0.00, 0.00)Client ID: MIL-K-S15

059724-0009-SA LAB ID:

Sampled: 30 MAR 98 Prepared: 06 APR 98 Dilution: 1.0 Received: 02 APR 98 SOIL Matrix: Analyzed: 23 APR 98 02 APR 98 Authorized:

HPLC-Y Instrument:

Parameter	Result	Qualifier	RL	MDL Units
Acenaphthene Acenaphthylene Anthracene Benzo(a)anthracene Benzo(b)fluoranthene Benzo(g,h,i)perylene Benzo(k)fluoranthene Chrysene Dibenz(a,h)anthracene Fluorene Fluoranthene Indeno(1,2,3-cd)pyrene Naphthalene Phenanthrene Pyrene	ND ND ND ND ND ND ND ND ND ND ND ND ND N		0.22 0.22 0.022 0.016 0.013 0.055 0.012 0.044 0.022 0.044 0.033 0.22 0.044 0.033	0.040 mg/kg 0.054 mg/kg 0.0029 mg/kg 0.0037 mg/kg 0.0024 mg/kg 0.0027 mg/kg 0.0033 mg/kg 0.0038 mg/kg 0.0032 mg/kg 0.0037 mg/kg 0.0037 mg/kg 0.0030 mg/kg 0.0030 mg/kg 0.0030 mg/kg 0.0031 mg/kg 0.0030 mg/kg
Surrogate		Recovery	. Acc	ceptable Range

100

*

22 - 167

Percent moisture is 9.0%. All results and limits are reported on a dry weight basis.

ND = Not Detected

Terphenyl-d14

Reported By: Blake Besser Approved By: Audrey Cornell



AFCEE Polynuclear Aromatic Hydrocarbons, HPLC Method 8310

Client Name: Parsons Engineering Science

Client ID: MIL-V-S26 (0.00, 0.00)

059724-0010-SA LAB ID:

SOIL Sampled: 29 MAR 98 Received: 02 APR 98 Matrix: Prepared: 06 APR 98 02 APR 98 Analyzed: 23 APR 98 Authorized:

Instrument.

Instrument: HPLC-Y	Dilution: 1.0	·
Parameter	Result Qualifier	RL MDL Units
Acenaphthene Acenaphthylene Anthracene Benzo(a)anthracene Benzo(b)fluoranthene Benzo(g,h,i)perylene Benzo(k)fluoranthene Chrysene Dibenz(a,h)anthracene Fluorene Fluoranthene Indeno(1,2,3-cd)pyrene Naphthalene Phenanthrene Pyrene	ND ND 0.074 M 0.069 M 0.11 M 0.063 M 0.037 M 0.11 M ND 0.015 Jd 0.17 M ND ND ND 0.15 d 0.18 d	0.24 0.044 mg/kg 0.24 0.059 mg/kg 0.024 0.0031 mg/kg 0.024 0.0041 mg/kg 0.018 0.0027 mg/kg 0.015 0.0030 mg/kg 0.061 0.0036 mg/kg 0.013 0.0042 mg/kg 0.048 0.0024 mg/kg 0.048 0.0024 mg/kg 0.048 0.0077 mg/kg 0.048 0.0079 mg/kg 0.048 0.0029 mg/kg 0.048 0.0036 mg/kg 0.048 0.0036 mg/kg 0.048 0.0036 mg/kg 0.048 0.0056 mg/kg 0.048 0.0067 mg/kg 0.048 0.0031 mg/kg
Surrogate	Recovery	Acceptable Range

100 Terphenyl-d14 * 22 - 167

Percent moisture is 17.4%. All results and limits are reported on a dry weight basis.

d = See Preferred Result on Other Column

J = Result is detected below the reporting limit or is an estimated concentration. M = Preferred Result

ND = Not Detected

Reported By: Blake Besser

Approved By: Audrey Cornell



AFCEE Polynuclear Aromatic Hydrocarbons, HPLC (Confirmation) Method 8310

Client Name: Parsons Engineering Science

Client ID: MIL-V-S26 (0.00,0.00)

LAB ID: 059724-0010-SA

Matrix: SOIL Sampled: 29 MAR 98 Received: 02 APR 98 Authorized: 02 APR 98 Prepared: 06 APR 98 Analyzed: 23 APR 98

Instrument: HPLC-Y Dilution: 1.0

This is the Lot	Dilacion. 1.0		
Parameter	Result Qualifier	RL	MDL Units
Acenaphthene Acenaphthylene Anthracene Benzo(a)anthracene Benzo(b)fluoranthene Benzo(g,h,i)perylene Benzo(k)fluoranthene Chrysene Dibenz(a,h)anthracene Fluorene Fluoranthene Indeno(1,2,3-cd)pyrene Naphthalene Phenanthrene Pyrene	ND ND 0.086 d 0.091 d 0.14 d 0.069 d 0.059 d 0.14 d ND 0.015 JM 0.22 d ND ND 0.11 M 0.16 M	0.24 0.24 0.024 0.018 0.015 0.061 0.013 0.048 0.024 0.048 0.048 0.036 0.24 0.048	0.044 mg/kg 0.059 mg/kg 0.0031 mg/kg 0.0041 mg/kg 0.0027 mg/kg 0.0030 mg/kg 0.0036 mg/kg 0.0042 mg/kg 0.0024 mg/kg 0.0041 mg/kg 0.0041 mg/kg 0.0077 mg/kg 0.0029 mg/kg 0.0036 mg/kg 0.0036 mg/kg 0.0056 mg/kg 0.0067 mg/kg 0.0031 mg/kg
Surrogate	Recovery	Acc	ceptable Range

96

*

22 - 167

Percent moisture is 17.4%. All results and limits are reported on a dry weight basis.

d = See Preferred Result on Other Column

J = Result is detected below the reporting limit or is an estimated concentration.

M = Preferred Result

ND = Not Detected

Terphenyl-d14

Reported By: Blake Besser

Approved By: Audrey Cornell



Method FL-PRO - TPH (C8-C40) Method FL-PRO

Client Name:

Parsons Engineering Science MIL-B-S25 Client ID: (0.00, 0.00)

059724-0008-SA LAB ID:

Sampled: 29 MAR 98 Prepared: 08 APR 98 Dilution: 1.0 SOIL Matrix: Received: 02 APR 98 02 APR 98 Authorized: Analyzed: 14 APR 98

Instrument: GCFID-I

Result Qualifier Parameter RL MDL Units 4.4 J 9.9 TPH (C8-C40) 11 mg/kg Surrogate Recovery Acceptable Range

o-Terphenyl Nonatriacontane 99 45 * * 22 - 166 10 - 192

J = Result is detected below the reporting limit or is an estimated concentration.

Reported By: Quanterra-Tampa Approved By: Cynthia Prentice



Method FL-PRO - TPH (C8-C40) Method FL-PRO

Client Name: Parsons Engineering Science

Client ID: MIL-K-S15 (0.00,0.00)

LAB ID: 059724-0009-SA

Matrix: SOIL Sampled: 30 MAR 98 Received: 02 APR 98 Authorized: 02 APR 98 Prepared: 08 APR 98 Analyzed: 14 APR 98

Instrument: GCFID-I Dilution: 1.0

Result Qualifier RL MDL Parameter Units 6.5 J 9.9 11 mg/kg TPH (C8-C40) Acceptable Range Recovery Surrogate **%** 22 - 166 10 - 192 o-Terphenyl 59 Nonatriacontane

J = Result is detected below the reporting limit or is an estimated concentration.

Reported By: Quanterra-Tampa Approved By: Cynthia Prentice



Method FL-PRO - TPH (C8-C40) Method FL-PRO

Client Name:

Parsons Engineering Science MIL-V-S26 Client ID: (0.00, 0.00)

059724-0010-SA LAB ID:

Sampled: 29 MAR 98 Prepared: 08 APR 98 Dilution: 1.0 Received: 02 APR 98 Analyzed: 14 APR 98 SOIL Matrix: Authorized: 02 APR 98

Instrument: GCFID-I

Parameter Result Qualifier **RL** MDL Units 4.7 J TPH (C8-C40) 11 9.9 mg/kg Surrogate Recovery Acceptable Range o-Terphenyl Nonatriacontane 96 43 * * 22 - 166 10 - 192

J = Result is detected below the reporting limit or is an estimated concentration.

Reported By: Quanterra-Tampa Approved By: Cynthia Prentice



Client Name: Client ID:

Parsons Engineering Science MIL-V-S8 059724-0027-SA

(0.00,0.00)

LAB ID: Matrix:

Sampled: 29 MAR 98 Prepared: See Below

Received: 02 APR 98 Analyzed: See Below

Authorized:

SOIL 02 APR 98

Parameter	Result,	Qual Dil	MDL	Rep L	im Units	Method	Prepared Date	Date
Total Organic Carbon Total Organic	ND	1.0	550	2000	mg/kg	9060	09 APR 98	13 APR 98
Carbon	ND	1.0	550	2000	mg/kg	9060	09 APR 98	13 APR 98
Total Organic Carbon Total Organic	ND	1.0	550	2000	mg/kg	9060	09 APR 98	13 APR 98
Carbon	ND	1.0	550	2000	mg/kg	9060	09 APR 98	13 APR 98

ND = Not Detected

Reported By: Patty Jungk



SAMPLE DESCRIPTION INFORMATION for Parsons Engineering Science

Lab ID	Client ID	Matrix	Sampled Date Time	Received Date
059619-0001-SA 059619-0002-SA 059619-0003-SA 059619-0004-SA 059619-0005-SA 059619-0006-SA 059619-0007-SA 059619-0008-TB	BX-MW-C BX-MW-07 BX-MW-D BX-MP-1 BX-MP-2 Mil-EA-04 Mil-EA-40 TRIP BLANK	GRND-H20 GRND-H20 GRND-H20 GRND-H20 GRND-H20 GRND-H20 GRND-H20 WATER-QA	25 MAR 98 16:0 26 MAR 98 06:3 26 MAR 98 07:3 26 MAR 98 13:3 26 MAR 98 12:3 26 MAR 98 14:4 26 MAR 98 15:0 26 MAR 98	0 27 MAR 98 0 27 MAR 98 0 27 MAR 98 0 27 MAR 98 5 27 MAR 98

A

Quanterra

Shain of Custody Record

QUA-1124 0797							
client Parsons		Project Manager	John	John Hicks	80/25/26/08	Chain of Custody Number 5304	304
1700 Broadway Suitchoo		Telephone Number (Area Code)/Fax Number	83188	ax Number	Lab Number	Page of	
M W		Chally Nagel	Vacel 12	1 Eller Lativier	Analysi: (Attach list if more space reneeded)		
		Carrier/Waybill Number	Ь		אקרי אקרי ממן	Special Instructions	/ 3
Contractip)Irchase Order/Quote No.			Matrix	Containers & Preservatives	1 0 % 1 % 1 % 1 % 1 % 1 % 1 % 1 % 1 % 1	Conditions of Receipt	ipt
Sample I.D. No. and Description (Containers for each sample may be combined on one line)	Date	ancentry BUIL	ios ios	NªOH SUPCH NªOH HCI HNO2 HSON	PAI TEN TEN TEN TO TEN TO TEN		
	3/25/98	T400 V			\ \ \		
	3/24/98	0630 V			\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\		
-		0730					
&BX-MP-1	3/20/98	1330 1					İ
65 BX-MP-2 B	1/26/98	1230 ~			777777		
Mi1-EA-04 5	3/20/98	1445		1	127777		
01 Mil- EA-40 B	3/26/98	1500			77777		
& Trip Blank	-						
Possible Hazard Identification Non-Hazard	J Potson B	Samp Unknown	Sample Disposal Return To Client	Disposal By Lab	(A lee may be a Months known than 3 m.	(A lee may be assessed if samples are retained forget than 3 months!	
12 [7 21 Dave	•		10			
184 O 1940		Dale 100	Time 1	1. Received By	1/V	Date / Time	1
man last		J/0/0/18		1 mary	1 () in ling	18/1/6	2
2. neuropusned by		Darie	l MM6	& Hecany d By		DAIs / Time	
3. Relinquished By		Date	Типе	3. Received By		. Типе Типе	
Comments							

DISTRIBUTION: WHITE - Stave with the Samule: CANARY - Returned to Client with Remort - PINK - Field Copy

Sheets Analyte The "x" Samples

Please hold the ren	BTEX	IPAHs	ED8	Mortes		1. j	
3/25/1998 COC#1			1220	Methane	TRPH		7
BX-GWRW-6	x	IX	lx				—
BX-MW-4	×	lx	- ^	+			\exists /
BX-GWRW-4		12					7/
BX-MW-2		-					1/59602
BX-MW-20							7/3
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Eller - based on the number, please all 2 BTEX and I EDB to both order it purish. Also, much the set on 2 for present brakeye.

Puk,

Page 2



SAMPLE DESCRIPTION INFORMATION for Parsons Engineering Science

				San	np1e	ed	Red	ceive	ed
Lab ID	Client ID	Matrix	D	ate		Time	[Date	
059638-0001-SA 059638-0002-SA 059638-0003-SA 059638-0004-SA 059638-0005-TB	Mil-EA-10 Mil-EA-5 Mil-EA-8	GRND - H20 GRND - H20 GRND - H20 GRND - H20 WATER - QA	27 27 27	MAR MAR MAR	98 98 98	08:30 09:30 11:30	28 28 28	MAR MAR MAR	98 98 98

@vanterra

Custody Record

Chain of

Chain of Custody Number 12305 1840 Special Instructions/ Conditions of Receipt ó Tme 182/2 2/8/ Analysis (Argentist if more space inceded) Date 3/27/98 <u>ک</u> ک ☐ Disposel By Lab ☐ Archive For OIERMS Telephone Number (Area Code)/Fax Number
303 831 8100
Site Contact
Lind 11 1098 Ellen La Kirle
Carrier/Waybiti Number QC Requirements (Specify) HOPN HOPN Containers & Preservatives Reselved By 1. Received By 2. Reported By ЮН Project Manager John Hicks SONH POSZH 14B Skin Inflant Poison B Unknown | Helum To Client Sample Disposal Matrix <u></u> Time 7005 3/27/08 000 0930 0830 1130 Other Time Date 3/21/98 3/27/98 3/27/98 3/21/98 ☐ 21 Days Swite 900 20000 ZPO Date 14 Days Sample I.D. No. and Description (Containers for each sample may be combined on one line) 7 Days 1700 Broadway Project Name Eglin AFB Contract Purchas Edden Quote No. | Flammable Client Parsons 1 48 Hours Jemp Blank Possible Hazard Identification M:1-EA-5 M:1-EA-8 M:1-EA-10 Tum Around Time Required SETrip Blank BX-MM-4 Dever 2. Relinquished By 3. Relinquished By Non-Hazard 1. Relinguished By QUA-4124 0797 24 Hours Comments Address



SAMPLE DESCRIPTION INFORMATION for Parsons Engineering Science

			Sampled	Received
Lab ID	Client ID	Matrix	Date Ti	me Date
059724-0001-SA 059724-0003-SA 059724-0003-SA 059724-0004-SA 059724-0005-SA 059724-0006-SA 059724-0008-SA 059724-0010-SA 059724-0010-SA 059724-0010-SD 059724-0011-SA 059724-0011-SA 059724-0013-SA 059724-0013-SA 059724-0015-SA 059724-0015-SA 059724-0016-SA 059724-0018-SA 059724-0018-SA 059724-0019-SA 059724-0020-SA 059724-0021-SA 059724-0021-SA	BXSB07S4.5 BX-SB07-S7 BX-SB08-S5 BX-SB09-S8 BX-SB11-S5 BX-SB12-S7 BX-SB14-S7 MIL-B-S25 MIL-V-S26 MIL-V-S26 MIL-V-S26 BX-MP02-S3 BX-MP02-S3 BX-MP02-S5 BX-VMP2-S7 BX-VEW1-S3 BX-VEW1-S3 BX-VEW1-S7 BXVEW2S5.5 BX-SB06-S3 BX-SB06-S4 BX-SB06-S7 BX-SB01-S7 BX-SB01-S7 BX-SB03-S7 MIL-K-S8 MIL-V-S8 TRIP BLANK	SOIL SOIL SOIL SOIL SOIL SOIL SOIL SOIL	28 MAR 98 16 28 MAR 98 16 28 MAR 98 16 29 MAR 98 00 30 MAR 98 00 30 MAR 98 00 29 MAR 98 00 29 MAR 98 00 29 MAR 98 00 29 MAR 98 00 29 MAR 98 00 26 MAR 98 10 26 MAR 98 10 28 MAR 98 07 28 MAR 98 07 28 MAR 98 07 28 MAR 98 08 28 MAR 98 08 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 09 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 15 28 MAR 98 10 29 MAR 98 00	:00 02 APR 98 :00 02 APR 98 :30 02 APR 98 :45 02 APR 98 :00 02 APR 98 :00 02 APR 98 :00 02 APR 98 :00 02 APR 98 :00 02 APR 98 :00 02 APR 98 :00 02 APR 98 :00 02 APR 98 :00 02 APR 98 :15 02 APR 98 :15 02 APR 98 :25 02 APR 98 :27 02 APR 98 :28 02 APR 98 :28 02 APR 98 :29 02 APR 98 :20 02 APR 98
059724-0029-RB	RINSEATE	AQUEOUS	31 MAR 98 00:	00 02 APR 98

Chain of Custody Record

59724

Wuanterra

QUA-1124 0797						
Fluxons	Project Manager	Project Manager Hicks	.0		3/31/98	Chain of Custody Numbr 02307
Broadway &		(303) 831 - 8100	Number		Lab Number	S , C
Denver 26 30290		Naged St	Sig Contact Nogel Lab Contact a Pilling		Analysis (Attach list If more space is needed)	
Sin AFB		umber		3871		Section 1
ContractPurchase Orber/Quote No.		Matrix	Containers & Preservatives	112 3 W + 2		Special instructions/ Conditions of Receipt
Sample I.D. No. and Description (Containers for each sample may be combined on one line)	Time	saudun sos spes	HOPN HOPN HOPN HOSONH	STS HAS IRT		
01 BX-SBOT-54,5 3/28/98	198 16000	X		XXX		
82 BX - SB07 - S7 3/28/98	000 1000	X		XXX		
63.BX-5B08-55 B/28/98		X		XXX		
4.BX - 5BO9 - 58 3/29/48	18 0745	X		XXX		
05-BX-5B11-S5 5/29/98	88	×		×××		
66 BX-SB12-ST 3/30/98	d8	×		XXX		
67BX-5BH-S7 3/30/98	18,	×		XXX		
08 Mil-B-525 3/29/98	198	X		XXX		
09Mil-K-515 3/30/98	188	X		XXX		
10 Mil-V-Sa6, 3/29/98	186	_ ×		XXX		
10m3 Mil - V-Sale/MS 3/29/	84/	×		XXX		
1030 Mil - V-SOV/MSD 3/29/48	/48	×		X		
Possible Hazard Identification Non-Hazard	Sample	Sample Disposal Return To Client	Disposal By Lab Arction For	retitus For	(A lee may be	(A fee may be assessed if samples are relained
Required T Days 14 D	Other		OC Requirements (Specify)		ì	(control
and Mage	186/1/h	Time 1530	1. Received By	K	17 P.	Date Time
2. Relinquished By	Date	Time	2. Received By		1	Date
3. Reinquished By	Date	Time	3. Received By			Date Time
Comments						

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132

Chain of Custody Record



DUA-4124 0797						
Client	Project Manager	John Hicks	Ş		13/31/98	Chain of Custody Number 5306
Su.		18 1	16)/Fax Number 8100			Page of 3
3	80290 Sile Cont.	Cindu Nage	Ellen a Rivieve		Analysis (Attach list if more space is needed)	
Project Name Eqiin AFIS	Carrier/W	1		38T		
Contrack/Purchase Orden/Quote No.		Matrix	Containers & Preservatives	# W+		Special instructions/ Conditions of Receipt
Sample I.D. No. and Description (Containers for each sample may be combined on one line)	Date Time	pos pos prourby	HOPN HOPN DH SONH POSZH	TKF PALL STEX		
1 BX-MP02-53	3/4/98 1015	×		 X X X		
3 BX-MP02-35	3/26/98 1015	×		XXX		
SK-VMP2-53	3/25/98 0725	×		XXX		
4 BX-VMP2-SS	3/28/98 0725	X		×××		
FX-VMPa-57	3/28/98 0725	X		×××		
BX-VEWI-S3	3/28/98 10815	X		×		
7 BX -VEWI-S5	3/28/98 0815	Х		XXX		
8 BX - VEWI-ST	3/28/98 0815	X		XXX		
9-BX-VEW 2-55.5	3/28/98 10930	X		XXX		
WBX-SBOG-S3	3/28/98 1500	.×		××		
1 BX - SB04 - SH	3/28/98 1500	.×		XXX		
ST	3/28/98 11500	×		XX		
Possible Hazard Identification		Sample Disposal			(A lee may be	second if samples are relatived
15	Unknown	☐ Return To Client	U Disposal By Lab	☐ Archive For	Months longer than 3 m	longer than 3 months)
24 Hours 48 Hours 7 Days 14 Days	21 Deys Other		Control of the contro			
1. Relinquished By Made Magel	1/4	18/1ms	1. Received By	1. H	2,:1	LLJ-OK TIME
2. Reinquished By	DAIR	Time	2. Received By	1	howard	Date Time
3. Relinquished By	Date	Time	3. Received By			Date
Comments	-					

DISTRIBUTION: WHITF . Slays with the Samula: CANARY - Raturned to Cliant with Report PINK . First Cony

Chain of Custody Record

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QUA-1124 0797							
Parsons		Project Manager	ngo, John	John Hicks		80/18/2	Chain of Custody Number 2308
1700 Broadway	Switchoo	Telephone (303	Telephone Number (Area Codo)/Fax Number (303) 831 – 8100	3 OD		Lab Number	3,3
VW Sale	26 COS 40 80240	Sine Contact	Sin Contact	Lab Contact Lab Contact		Analysis (Attach list if more space is needed)	5
Project Name Eglin AFTS		Carrier/Wd/bill Number	į.		3877		
Contract/Purchasb_Opdar/Quote No.			Matrix	Containers & Preservatives	THE S		Special Instructions/ Conditions of Receipt
Sample I.D. No. and Description (Containers for each sample may be combined on one line)	Date	Тітв	pos pos snoenby	HOPH NEOH HCI HOSZH HSZOY NUDURE	PAH PAH AT TO		
25 BX-5B01-S7	3/28/98	0750	×		>		
24 BX-5802-ST	3/28/98	아이	×		×		
7	3/28/48	1120	×		×		
26 Mil-K-S8	3/30/48		×		>		
27 Mil - V - S8	3/50/68		×		>		
Mrip Blank	3/31/88		 	×	XXX		
Schalake	3/31/98		×				
n mmable 🔲 Skin Inflant	Polson B	Sal Cinknown	Sample Disposal Return To Client	Disposal By Lab Archive For	Archive For	(A lee may be ass	(A lee may be assessed if samples are relained
Required 7 Days 140	0 21 08	4 .			<i>(</i> / ₄)		fem
1. Relinquished By/ MMM-TMBO	00	5/1/4%	8/1550	1. Received By	Kust R	المراز الم	Date Time
		Dile	Time	2. Received By			
3. Relinquished By		Date	Time	3. Received By			Date
Comments							

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3/25/98 1	PROJECT NAME EGIN AFB, F	NOTEBOOK NO.
	Report of field analysis of	groundwater samples at (1th St. Service Station)
	Well Analyte Method	Concentration(my/L) Sample Time
	GWRW-6 ferrousiron HACH Colori DR/700 V.	metric 0.00 mg/L 0730
	GWRW-6 surfate HACH Cold DR/700 V	rimetric 131 mg/L 0730
	MW-4 ferrous ivon sulfate	0.03 mg/L 0900 71 mg/L 0900
	GWRW-4 ferrousian	0.08 mg/L 0800
	sufate	161 mg/L 0800 0.13 mg/L 1200
	MW-2 ferrous uon	27 mg/L
	MW-2 (dup) ferrous ion	- 0.12 mg/L 1200 36 mg/L
	MW-1 Jenous iner	
	MW-C Serrous iron	0.01 mg/L 1600 45 mg/L
	sulpte	TO MG/L
		·
	SIGNATURE Cindy Nag	DATE MACK 25 1

3/26/98 2	PROJECT NAME Eglin AFB	NOTEBOOK NO
	Record of field analysis of ground was	er samples.
	Using HACH coloringter to analy	
	fermus iron DE 1000 Mayor	Ze tor
	ferrous ivon (DR/700 Method Sulfate (DR/700 Method	150.01) and
1	Bx Service Station	75.088
	BX Service Station 0630 MW-7 ferrous iron	m 01 mm/1
	sulfate	0.01 mg/L 155 mg/L
		1976
	0730 MW-D ferreus iron	0.07 mg/L
	sulpte	89 mg/L
		<u> </u>
	1230 MP-2 Jerrous iran	
	1230 MP-2 ferrous iron	0.55 mg/L 152 mg/L
	Daulfate	132 mg/L
	MP-2 duplicate ferrousiron	^ =/- /·
	sulfate	0.56 mg/L
	sugar	150 mg/L
	1330 MP-1 Jerrous iron	000
	sulpate	0.00 mg/L 236 mg/L
		mg/c
	Military Gas Station	
	1445 EA-4 Jehnous 1000	
		0.40 mg/L
) sulfate	67 mg/L
	FA-4 due terrous Hon	
	EA-4 dup ferrous von	0.40 mg/
	**************************************	- 89 mg/L
	SIGNATURE A 11 AA	
	READ AND UNDERSTOOD	_ DATE 3/26 1998
	LILAD VIAD GIADERS LOGD	_ DATE 719

3/27/08 3	PROJECT	NAME agun	Arb, Stonda	NOTEBOOK NO
	Record	of field as	rolysis & ground	water Aamples
	110.	e Hary Non	Dain iti I auan	in you
	MAIN	Jenous is	eriniting to anal ion (DR/700 Method tran	200 50.01)
	and	2 sulfate	(DR/700 Method	45.000
	Militar	y bas Sta	tion	<i></i>
	0830	EA-10	Jerrous iron	0.00 mg/L
			sulfate	76 mg/L
	1030	EA-5	1455 154	<u> </u>
	0930	<u> </u>	Jerrous tran	0.01 mg/L 120 mg/L
			<u> </u>	
	1130	EA-8	Jerrous iron	0.00 mg/L
			sulfate	109 mg/L

	-			
	SIGNATUF	RE Hinds	Magol	DATE 3/27 19
		UNDERSTOOD		DATE19
				•

AIR TOXICS LTD.

SAMPLE NAME: Mil SG1

ID#: 9804028-04A

EPA Method TO-3 GC/PID/FID

File Name*	
	Date of Collection: 4/ 1/98
Dil Factor	Date of Analysis: 4/15/98

Compound	Rpt. Limit (ppmv)	Rpt. Limit (uG/L)	Amount (ppmv)	Amount (uG/L)
Benzene	0.020	0.064	Not Detected	Not Detected
Toluene	0.020	0.075	0.026	0.099
Ethyl Benzene	0.020	0.087	Not Detected	Not Detected
Total Xylenes	0.020	0.087	0.075	0.33
TPH (C5+ Hydrocarbons) ref. to Gasoline	0.20	0.82	1.5	6.2
C2-C4 Hydrocarbons ref. to Gasoline	0.20	0.36	Not Detected	Not Detected

Container Type: 1 Liter Summa Canister

Surrogates		Method
	% Recovery	Limits
Fluorobenzene (PID)	116	50-150
Fluorobenzene (FID)	116	50-150

3/31/98 4	PROJECT NAME <u>Eglin AFB</u> , <u>Florida</u> NOTEBOOK NO
<u> </u>	Record of field analysis for groundwater samples.
<u>-</u>	Using HACH colorineter
	Method DR/700 50.01 ferrous iron
	Hainer HACH colorimeter Method DR/700 50.01 ferrous iron Method DR/700 45.000 sulfate
ri,	TRY Service Station SB-15/MD3
	Bl Service Station SB-15/MP3 ferrous iron 3.2 mg/L suffate 340 mg/L
	sulpte 340 mg/L
	7
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19	SIGNATURE [moly Magel DATE 3/3/ 1998
វ	READ AND UNDERSTOOD DATE 3/3/ 19 98

APPENDIX B DATA FROM PREVIOUS INVESTIGATIONS

SUMMARY OF DRILLING AND WELL CONSTRUCTION L.. FOR MILITARY GAS STATION ON MAIN BASE, EGLIN AFB, VALPARAISO, OKALOOSA COUNTY, FLORIDA. TABLE,

	, in the	Top of		E	Screen/Casi	Screen/Casing Interval ^{®)}	1		
Drilling Location	Drill Installation Date	Casing Elevation ⁽⁴⁾ (ft MSL)	Ground Elevation (ft MSL)	Total Depth . (ft)	Screen (ft)	Riser (fi)	Sandpack Interval (fi)	Benionile Seal (fi)	Grout (ft)
EA-1	8-24-92	54.00	54.28	45	45.0-30.0	30.0-0	45.0-28.0	28.0-26.0	26.0-0
EA-2	8-24-92	54.08	54.34	45	45.0-30.0	30.0-0	45.0-28.0	28.0-26.0	26.0-0
EA-3	8-25-92	54.32	54.57	45	45.0-30.0	30.0-0	45.0-28.0	28.0-26.0	26.0-0
EA-4	8-26-92	55.47	55.82	45	45.0-30.0	30.0-0	45.0-28.0	28.0-26.0	26.0-0
EA-5	8-27-92	55.24	55.40	45	45.0-30.0	30.0-0	45.0-28.0	28.0-26.0	26.0-0
EA-6	8-28-92	55.85	56.12	45	45.0-30.0	30.0-0	45.0-28.0	28.0-26.0	26.0-0
EA-7	8-28-92	53.94	54.19	45	45.0-30.0	30.0-0	45.0-28.0	28.0-26.0	26.0-0
EA-8	8-31-92	56.04	56.30	45	45.0-30.0	30.0-0	45.0-28.0	28.0-26.0	26.0-0
EA-9	9-1-92	55.43	55.74	59	57.0-54.5	54.5-0	59.0-53.5	53.5-52.0	52.0-0
EA-10	6-14-93	54.72	55.07	45	45.0-30.0	30.0-0	45.0-28.0	28.0-26.0	26.0-0
EA-11	6-14-93	53.58	53.91	45	45.0-30.0	30.0-0	45.0-28.0	28.0-26.0	26.0-0
≺	8-19-92	Y.	53.9	42	ı	ı	ı	i	42.0-0
æ	8-18-92	Y _N	54.2	42	ı	1	:	:	42.0-0
ပ	8-20-92	Y.	53.9	45	1	ı	ı	ı	42.0-0
D	8-20-92	NA	54.3	42	1	ł	ı		45.0-0
11	8-24-92	٧٧	53.6	42	1	ŀ	ı	ı	45.0-0
•	8-26-92	Ϋ́N	55.2	42	1		1		42.0-0
×	8-26-92	Y _N	55.0	42	1	ı	1	ı	45.0-0
1	8-26-92	NA	55.1	42	1	1	ı	ı	42.0-0
0	6-12-93	NA	54.99	36	1	1	1	ı	36.0-0
æ	6-12-93	NA	54.97	36	1	1	1	ı	36.0-0
S	6-15-93	VV	55.82	36	1	ı	i	ı	36.0-0

⁽a) = Elevations of EA-10, EA-11, borings Q through W, AAS-1, SVE-1, and SVE-2 surveyed by George, Nielsen, and Tooke, P.A. of Crestview, Florida on 14 July 1993. Elevations of EA-1 through EA-9 and borings A through L surveyed by Dowco Land Surveying of Ft. Walton Beach, Florida on 8 September 1992.

(b) = All monitoring wells constructed with 2-in. diameter, Schedule 40 PVC screen and casing.

(c) = AAS-1 has a PVC sump from 60.0 to 57.5 ft below ground surface.

) in C	Top of	Parion	Total	Screen/Casing Interval ⁽⁶⁾	g Interval ⁽⁶⁾	Candonal	Descrite	
Drilling Location	Installation Date	Elevation ^(a) (ft MSL)	Elevation (ft MSL)	Depth (ft)	· Screen (fi)	Riser (ft)	Interval (ft)	Seal (ft)	Grout (ft)
۲	6-15-93	NA NA	55.90	36	:	ł	ı	ı	36.0-0
Ω	6-12-93	YN	55.63	36	1	ı		1	36.0-0
>	6-16-93	ΥN	56.16	36	ı	1	:	I	36.0-0
*	6-17-93	YY	56.24	36	ı	1	·	1	36.0-0
AAS-1	5-13-93	55.21	55.46	19	57.5-55.0(4)	55.0-0	60.0-49.0	49.0-47.0	47.0-0
SVE-1	5-11-93	55.34	55.58	32	32.0-7.0	7.0-0	32.0-5.0	5.0-3.0	3.0-0
SVE-2	5-11-93	54.54	55.22	32	32.0-7.0	7.0-0	32.0-5.0	5.0-3.0	3.0-0

TABLE 2

PUBLIC GROUNDWATER SUPPLY WELLS AT THE MAIN BASE, EGLIN AIR FORCE BASE, VALPARAISO, OKALOOSA COUNTY, FLORIDA⁽⁴⁾.

E	g water						
USE	Drinking water	Drinking water	Drinking water	Drinking water	Drinking water	Drinking water	Drinking water
AQUIFER	Floridan						
PUMP DEPTH (ft.)	168	190	150	165	150	180	unknown
GPM	300	1000	700	1000	950	. 400	350
DEPTH (FT.)	575	209	652	550	642	290	702
SIZE (in.)	10	12	12	12	16	9	∞
WELL NO.	yand	7	ĸ	4	5	9	61

Information from records on file at Base Civil Engineering office, Eglin AFB.

TABLE 2

PUBLIC GROUNDWATER SUPPLY WELLS AT THE MAIN BASE, EGLIN AIR FORCE BASE, VALPARAISO, OKALOOSA COUNTY, FLORIDA®.

AQUIFER USE	Floridan Drinking water	Floridan Drinking water	Floridan Drinking water	Floridan Drinking water	Floridan Drinking water	Floridan Drinking water	Floridan Drinking water	
PUMP DEPTH //	168	190	150	165	150	180	unknown	
ВРМ	300	1000	700	1000	950	400	350	
DEPTH (FT.)	575	209	652	550	642	290	702	
SIZE (in.)	10	12	12	12	16	9	∞	
WELL NO.		2	т	4	2	9	61	

Information from records on file at Base Civil Engineering office, Eglin AFB.

TABLE 4 SUMMARY OF GAUGING DATA FOR MILITARY GAS STATION ON MAIN BASE, EGLIN AFB, VALPARAISO, OKALOOSA COUNTY, FLORIDA.

Date Number Elevation To LPH Elevation (feet) Water Elev Thickness (feet)		Well	Casing	Depth	LPH	Depth To	Corrected	LPH
01-Sep-92 EA-1 *N/A N/A N/A 34.50 N/A N/A 09-Sep-92 EA-1 54.00 N/A N/A 34.26 19.74 N/A N/A 13-Sep-92 EA-1 54.00 N/A N/A 34.26 19.74 N/A 13-Sep-92 EA-1 54.00 N/A N/A 34.26 19.74 N/A 13-Sep-92 EA-1 54.00 N/A N/A 34.31 19.69 N/A 22.Jan-93 EA-1 54.00 N/A N/A 34.69 19.31 N/A 18-Jun-93 EA-1 54.00 N/A N/A 35.58 18.42 N/A 18-Jun-93 EA-1 54.00 N/A N/A 36.69 19.31 N/A 27-Jul-93 EA-1 54.00 N/A N/A 36.69 17.31 N/A 17-Aug-93 EA-1 54.00 N/A N/A 36.69 17.31 N/A 17-Aug-93 EA-1 54.00 N/A N/A 36.69 17.31 N/A 16-Sep-93 EA-1 54.00 N/A N/A 36.69 17.31 N/A 16-Sep-93 EA-1 54.00 N/A N/A 36.61 17.39 N/A 16-Sep-93 EA-1 54.00 N/A N/A 36.61 17.39 N/A 11-Feb-94 EA-1 54.00 N/A N/A 36.71 17.29 N/A 11-Feb-94 EA-1 54.00 N/A N/A 36.71 17.29 N/A 11-Feb-94 EA-1 54.00 N/A N/A 36.16 17.84 N/A 11-Beb-94 EA-1 54.00 N/A N/A 36.16 17.84 N/A 08-Apr-94 EA-1 54.00 N/A N/A 35.70 18.30 N/A 08-Apr-94 EA-1 54.00 N/A N/A 35.70 18.30 N/A 08-Apr-94 EA-1 54.00 N/A N/A 36.73 17.27 N/A 02-Jun-94 EA-1 54.00 N/A N/A 36.91 17.09 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-2 54.08 N/A N/A 31.59 22.41 N/A N/A 31.59 22.41 N/A N/A 31.59 22.41 N/A N/A 31.59 22.41 N/A N/A 31.59 22.41 N/A N/A 31.59 22.41 N/A N/A 31.59 22.41 N/A N/A 31.59 22.59.94 EA-2 54.08 N/A N/A 31.50 0 N/A N/A 31.50 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.99 EA-2 54.08 N/A N/A	Date			•	Elevation	•		
09-Sep-92 EA-1 54.00 N/A N/A 34.26 19.74 N/A 13-Sep-92 EA-1 54.00 N/A N/A 34.31 19.69 N/A 22-Jan-93 EA-1 54.00 N/A N/A 34.31 19.69 N/A 21-May-93 EA-1 54.00 N/A N/A 34.63 19.31 N/A 21-May-93 EA-1 54.00 N/A N/A 34.69 19.31 N/A 18-Jun-93 EA-1 54.00 N/A N/A 36.69 17.31 N/A 17-Jun-93 EA-1 54.00 N/A N/A 36.69 17.31 N/A 17-Jun-93 EA-1 54.00 N/A N/A 36.69 17.31 N/A 17-Jun-93 EA-1 54.00 N/A N/A 36.61 17.39 N/A 16-Sep-93 EA-1 54.00 N/A N/A 36.61 17.39 N/A 16-Sep-93 EA-1 54.00 N/A N/A 36.31 17.52 N/A 05-Oct-93 EA-1 54.00 N/A N/A 36.31 17.52 N/A 11-Feb-94 EA-1 54.00 N/A N/A 36.31 17.52 N/A 11-Feb-94 EA-1 54.00 N/A N/A 37.34 16.66 N/A 07-Mar-94 EA-1 54.00 N/A N/A 35.70 18.30 N/A 19-May-94 EA-1 54.00 N/A N/A 35.70 18.30 N/A 19-May-94 EA-1 54.00 N/A N/A 35.70 18.30 N/A 19-May-94 EA-1 54.00 N/A N/A 36.73 17.27 N/A 02-Jun-94 EA-1 54.00 N/A N/A 36.73 17.27 N/A 02-Jun-94 EA-1 54.00 N/A N/A 36.91 17.09 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-2 54.08 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-2 54.08 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-2 54.08 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50					(feet MSL)	(feet)		
09-Sep-92 EA-1 54.00 N/A N/A 34.26 19.74 N/A 13-Sep-92 EA-1 54.00 N/A N/A 34.31 19.69 N/A 22-Jan-93 EA-1 54.00 N/A N/A 34.31 19.69 N/A 21-May-93 EA-1 54.00 N/A N/A 34.63 19.31 N/A 21-May-93 EA-1 54.00 N/A N/A 34.69 19.31 N/A 18-Jun-93 EA-1 54.00 N/A N/A 36.69 17.31 N/A 17-Jun-93 EA-1 54.00 N/A N/A 36.69 17.31 N/A 17-Jun-93 EA-1 54.00 N/A N/A 36.69 17.31 N/A 17-Jun-93 EA-1 54.00 N/A N/A 36.61 17.39 N/A 16-Sep-93 EA-1 54.00 N/A N/A 36.61 17.39 N/A 16-Sep-93 EA-1 54.00 N/A N/A 36.31 17.52 N/A 05-Oct-93 EA-1 54.00 N/A N/A 36.31 17.52 N/A 11-Feb-94 EA-1 54.00 N/A N/A 36.31 17.52 N/A 11-Feb-94 EA-1 54.00 N/A N/A 37.34 16.66 N/A 07-Mar-94 EA-1 54.00 N/A N/A 35.70 18.30 N/A 19-May-94 EA-1 54.00 N/A N/A 35.70 18.30 N/A 19-May-94 EA-1 54.00 N/A N/A 35.70 18.30 N/A 19-May-94 EA-1 54.00 N/A N/A 36.73 17.27 N/A 02-Jun-94 EA-1 54.00 N/A N/A 36.73 17.27 N/A 02-Jun-94 EA-1 54.00 N/A N/A 36.91 17.09 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-2 54.08 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-2 54.08 N/A N/A 31.59 22.41 N/A 07-Jun-94 EA-2 54.08 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50 0 N/A N/A 31.50								
13.Sep-92 EA-1 54.00 N/A N/A 34.89 19.31 N/A 22-Jan-93 EA-1 54.00 N/A N/A 34.89 19.31 N/A 18-Jun-93 EA-1 54.00 N/A N/A 35.88 18.42 N/A 18-Jun-93 EA-1 54.00 N/A N/A 36.89 17.31 N/A 27-Jul-93 EA-1 54.00 N/A N/A 36.89 17.51 N/A 17-Aug-93 EA-1 54.00 N/A N/A 36.89 17.31 N/A 17-Aug-93 EA-1 54.00 N/A N/A 36.89 17.31 N/A 17-Aug-93 EA-1 54.00 N/A N/A 36.81 17.39 N/A 16-Sep-93 EA-1 54.00 N/A N/A 36.81 17.39 N/A 16-Sep-93 EA-1 54.00 N/A N/A 36.81 17.39 N/A 11-Feb-94 EA-1 54.00 N/A N/A 36.73 17.27 N/A 11-Feb-94 EA-1 54.00 N/A N/A 36.73 17.27 N/A 19-May-94 EA-1 54.00 N/A N/A 35.70 18.30 N/A 19-May-94 EA-1 54.00 N/A N/A 35.70 18.30 N/A 19-May-94 EA-1 54.00 N/A N/A 36.73 17.27 N/A 19-May-94 EA-1 54.00 N/A N/A 36.73 17.27 N/A 102-Jun-94 EA-1 54.00 N/A N/A 36.73 17.27 N/A 17-Jun-94 EA-1 54.00 N/A N/A 36.73 17.27 N/A 17-Jun-94 EA-1 54.00 N/A N/A 36.73 17.27 N/A 17-Jun-94 EA-1 54.00 N/A N/A 36.91 17.09 N/A 11-Aug-94 EA-1 54.00 N/A N/A 34.64 19.36 N/A 11-Aug-94 EA-1 54.00 N/A N/A 34.64 19.36 N/A 11-Aug-94 EA-1 54.00 N/A N/A 34.64 19.36 N/A 11-Aug-94 EA-1 54.00 N/A N/A 34.64 19.36 N/A 11-Aug-94 EA-1 54.00 N/A N/A 34.64 19.36 N/A 11-Aug-94 EA-1 54.00 N/A N/A 34.84 19.36 N/A 11-Aug-94 EA-1 54.00 N/A N/A 34.84 19.36 N/A 11-Aug-94 EA-1 54.00 N/A N/A 34.80 19.96 N/A 11-Aug-94 EA-2 54.08 N/A N/A 34.10 19.98 N/A 11-Aug-94 EA-2 54.08 N/A N/A 34.10 19.98 N/A 11-Aug-94 EA-2 54.08 N/A N/A 34.10 19.98 N/A 11-Aug-94 EA-2 54.08 N/A N/A 34.10 19.98 N/A 11-Aug-94 EA-2 54.08 N/A N/A 34.10 19.98 N/A 11-Aug-94 EA-2 54.08 N/A N/A 36.50 17.75 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.50 17.50 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.50 17.20 N/A 11-Aug-94 EA-2 54.08								
22-Jan-93 EA-1 54.00 N/A N/A 34.69 19.31 N/A 21-May-93 EA-1 54.00 N/A N/A 35.58 18.42 N/A 18-Jun-93 EA-1 54.00 N/A N/A 36.69 17.51 N/A 27-Jul-93 EA-1 54.00 N/A N/A 36.69 17.31 N/A 17-Aug-93 EA-1 54.00 N/A N/A 36.69 17.31 N/A 17-Aug-93 EA-1 54.00 N/A N/A 36.69 17.31 N/A 17-Aug-93 EA-1 54.00 N/A N/A 36.69 17.31 N/A 05-Oct-93 EA-1 54.00 N/A N/A 36.38 17.62 N/A 05-Oct-93 EA-1 54.00 N/A N/A 36.38 17.62 N/A 11-Feb-94 EA-1 54.00 N/A N/A 36.31 17.29 N/A 11-Feb-94 EA-1 54.00 N/A N/A 36.16 17.84 N/A 21-Mar-94 EA-1 54.00 N/A N/A 35.70 18.30 N/A 08-Apr-94 EA-1 54.00 N/A N/A 35.70 18.30 N/A 19-May-94 EA-1 54.00 N/A N/A 35.70 18.30 N/A 19-May-94 EA-1 54.00 N/A N/A 35.70 18.30 N/A 19-May-94 EA-1 54.00 N/A N/A 36.73 17.27 N/A 02-Jun-94 EA-1 54.00 N/A N/A 36.91 17.09 N/A 07-Jul-94 EA-1 54.00 N/A N/A 34.64 19.36 N/A 11-Aug-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 22-Sep-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 22-Sep-94 EA-1 54.00 N/A N/A 31.59 22.41 N/A 12-Oct-94 EA-1 54.00 N/A N/A 32.32 N/A N/A 31.99 22.41 N/A 12-Oct-94 EA-1 54.00 N/A N/A 34.16 19.92 N/A 12-Oct-94 EA-2 54.08 N/A N/A 34.16 19.92 N/A 13-Sep-92 EA-2 54.08 N/A N/A 34.16 19.92 N/A 13-Sep-92 EA-2 54.08 N/A N/A 36.33 17.75 N/A 13-Sep-92 EA-2 54.08 N/A N/A 36.33 17.75 N/A 13-Sep-92 EA-2 54.08 N/A N/A 36.50 17.78 N/A 13-Sep-92 EA-2 54.08 N/A N/A 36.50 17.78 N/A 13-Sep-93 EA-2 54.08 N/A N/A 36.50 17.78 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.50 17.78 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.50 17.79 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.50 17.79 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.50 17.79 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.50 17.79 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.50 17.79 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.50 17.79 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.50 17.79 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.50 17.79 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.50 17.50 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.50 17.50 N/A N/A 36.50 17.50 N/A N/A 36.50 17.50 N/A N/A 36.50 17	09-Sep-92	EA-1						
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18-Jun-93 EA-2 54.08 N/A N/A 36.33 17.75 N/A 27-Jul-93 EA-2 54.08 N/A N/A 36.52 17.56 N/A 17-Aug-93 EA-2 54.08 N/A N/A 36.47 17.61 N/A 16-Sep-93 EA-2 54.08 N/A N/A 36.30 17.78 N/A 05-Oct-93 EA-2 54.08 N/A N/A 36.30 17.78 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.53 17.55 N/A 11-Feb-94 EA-2 54.08 N/A N/A 37.18 16.90 N/A 07-Mar-94 EA-2 54.08 N/A N/A 35.56 18.52 N/A 08-Apr-94 EA-2 54.08 N/A N/A 35.76 18.32 N/A 19-May-94 EA-2 54.08 N/A N/A 36.54 17.54 N/A 07-Jun-94	22-Jan-93	EA-2						
27-Jul-93 EA-2 54.08 N/A N/A 36.52 17.56 N/A 17-Aug-93 EA-2 54.08 N/A N/A 36.47 17.61 N/A 16-Sep-93 EA-2 54.08 N/A N/A 36.30 17.78 N/A 05-Oct-93 EA-2 54.08 N/A N/A 36.53 17.55 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.53 17.55 N/A 11-Feb-94 EA-2 54.08 N/A N/A 37.18 16.90 N/A 07-Mar-94 EA-2 54.08 N/A N/A 35.56 18.52 N/A 21-Mar-94 EA-2 54.08 N/A N/A 35.56 18.52 N/A 08-Apr-94 EA-2 54.08 N/A N/A 35.76 18.32 N/A 19-May-94 EA-2 54.08 N/A N/A 36.54 17.54 N/A 02-Jun-94 EA-2 54.08 N/A N/A 36.86 17.22 N/A 07-Jun-94 EA-2 54.08 N/A N/A 36.86 17.22 N/A 07-Jun-94 EA-2 54.08 N/A N/A 36.70 17.38 N/A 07-Jun-94 EA-2 54.08 N/A N/A 36.18 17.90 N/A 07-Jul-94 EA-2 54.08 N/A N/A 36.18 17.90 N/A 11-Aug-94 EA-2 54.08 N/A N/A 34.71 19.37 N/A 11-Aug-94 EA-2 54.08 N/A N/A 32.63 21.45 N/A 12-Oct-94 EA-2 54.08 N/A N/A 32.63 21.45 N/A 12-Oct-94 EA-2 54.08 N/A N/A 34.36 19.96 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A	21-May-93	EA-2						
17-Aug-93 EA-2 54.08 N/A N/A 36.47 17.61 N/A 16-Sep-93 EA-2 54.08 N/A N/A 36.30 17.78 N/A 05-Oct-93 EA-2 54.08 N/A N/A 36.53 17.55 N/A 11-Feb-94 EA-2 54.08 N/A N/A 36.53 17.55 N/A 07-Mar-94 EA-2 54.08 N/A N/A 36.00 18.08 N/A 07-Mar-94 EA-2 54.08 N/A N/A 35.56 18.52 N/A 08-Apr-94 EA-2 54.08 N/A N/A 35.56 18.52 N/A 08-Apr-94 EA-2 54.08 N/A N/A 35.76 18.32 N/A 19-May-94 EA-2 54.08 N/A N/A 36.54 17.54 N/A 02-Jun-94 EA-2 54.08 N/A N/A 36.86 17.22 N/A 07-Jun-94 EA-2 54.08 N/A N/A 36.86 17.22 N/A 07-Jun-94 EA-2 54.08 N/A N/A 36.70 17.38 N/A 28-Jun-94 EA-2 54.08 N/A N/A 36.18 17.90 N/A 07-Jul-94 EA-2 54.08 N/A N/A 36.18 17.90 N/A 11-Aug-94 EA-2 54.08 N/A N/A 34.71 19.37 N/A 11-Aug-94 EA-2 54.08 N/A N/A 34.71 19.37 N/A 11-Aug-94 EA-2 54.08 N/A N/A 31.42 22.66 N/A 12-Oct-94 EA-2 54.08 N/A N/A 31.94 22.14 N/A 01-Sep-92 EA-3 54.08 N/A N/A 34.36 19.96 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A	18-Jun-93	EA-2						
16-Sep-93	27-Jul-93	EA-2						
05-Oct-93	17-Aug-93	EA-2						
11-Feb-94 EA-2 54.08 N/A N/A 37.18 16.90 N/A 07-Mar-94 EA-2 54.08 N/A N/A 36.00 18.08 N/A 21-Mar-94 EA-2 54.08 N/A N/A 35.56 18.52 N/A 08-Apr-94 EA-2 54.08 N/A N/A 35.76 18.32 N/A 19-May-94 EA-2 54.08 N/A N/A 36.54 17.54 N/A 02-Jun-94 EA-2 54.08 N/A N/A 36.54 17.22 N/A 07-Jun-94 EA-2 54.08 N/A N/A 36.86 17.22 N/A 07-Jun-94 EA-2 54.08 N/A N/A 36.70 17.38 N/A 28-Jun-94 EA-2 54.08 N/A N/A 36.18 17.90 N/A 07-Jul-94 EA-2 54.08 N/A N/A 34.71 19.37 N/A 11-Aug-94 EA-2 54.08 N/A N/A 34.71 19.37 N/A 11-Aug-94 EA-2 54.08 N/A N/A 31.42 22.66 N/A 22-Sep-94 EA-2 54.08 N/A N/A 31.42 22.66 N/A 12-Oct-94 EA-2 54.08 N/A N/A 31.94 22.14 N/A 01-Sep-92 EA-3 54.32 N/A N/A 34.36 19.96 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A	16-Sep-93							
07-Mar-94 EA-2 54.08 N/A N/A 36.00 18.08 N/A 21-Mar-94 EA-2 54.08 N/A N/A 35.56 18.52 N/A 08-Apr-94 EA-2 54.08 N/A N/A 35.76 18.32 N/A 19-May-94 EA-2 54.08 N/A N/A 36.54 17.54 N/A 02-Jun-94 EA-2 54.08 N/A N/A 36.86 17.22 N/A 07-Jun-94 EA-2 54.08 N/A N/A 36.70 17.38 N/A 28-Jun-94 EA-2 54.08 N/A N/A 36.18 17.90 N/A 07-Jul-94 EA-2 54.08 N/A N/A 34.71 19.37 N/A 11-Aug-94 EA-2 54.08 N/A N/A 31.42 22.66 N/A 12-Oct-94 EA-2 54.08 N/A N/A 31.94 22.14 N/A 01-Sep-92	05-Oct-93	EA-2	54.08					
21-Mar-94 EA-2 54.08 N/A N/A 35.56 18.52 N/A 08-Apr-94 EA-2 54.08 N/A N/A 35.76 18.32 N/A 19-May-94 EA-2 54.08 N/A N/A 36.54 17.54 N/A 02-Jun-94 EA-2 54.08 N/A N/A 36.86 17.22 N/A 07-Jun-94 EA-2 54.08 N/A N/A 36.70 17.38 N/A 28-Jun-94 EA-2 54.08 N/A N/A 36.18 17.90 N/A 07-Jul-94 EA-2 54.08 N/A N/A 36.18 17.90 N/A 07-Jul-94 EA-2 54.08 N/A N/A 34.71 19.37 N/A 11-Aug-94 EA-2 54.08 N/A N/A 31.42 22.66 N/A 22-Sep-94 EA-2 54.08 N/A N/A 31.42 22.66 N/A 12-Oct-94 EA-2 54.08 N/A N/A 31.94 22.14 N/A 01-Sep-92 EA-3 54.32 N/A N/A 34.36 19.96 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A	11-Feb-94	EA-2	54.08					
08-Apr-94 EA-2 54.08 N/A N/A 35.76 18.32 N/A 19-May-94 EA-2 54.08 N/A N/A 36.54 17.54 N/A 02-Jun-94 EA-2 54.08 N/A N/A 36.86 17.22 N/A 07-Jun-94 EA-2 54.08 N/A N/A 36.70 17.38 N/A 28-Jun-94 EA-2 54.08 N/A N/A 36.18 17.90 N/A 07-Jul-94 EA-2 54.08 N/A N/A 36.18 17.90 N/A 07-Jul-94 EA-2 54.08 N/A N/A 34.71 19.37 N/A 11-Aug-94 EA-2 54.08 N/A N/A 31.42 22.66 N/A 22-Sep-94 EA-2 54.08 N/A N/A 31.42 22.66 N/A 12-Oct-94 EA-2 54.08 N/A N/A 31.94 22.14 N/A 01-Sep-92 EA-3 54.08 N/A N/A 34.36 19.96 N/A 09-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A	07-Mar-94	EA-2	54.08					
19-May-94 EA-2 54.08 N/A N/A 36.54 17.54 N/A 02-Jun-94 EA-2 54.08 N/A N/A 36.86 17.22 N/A 07-Jun-94 EA-2 54.08 N/A N/A 36.70 17.38 N/A 28-Jun-94 EA-2 54.08 N/A N/A 36.18 17.90 N/A 07-Jul-94 EA-2 54.08 N/A N/A 36.18 17.90 N/A 07-Jul-94 EA-2 54.08 N/A N/A 34.71 19.37 N/A 11-Aug-94 EA-2 54.08 N/A N/A 31.42 22.66 N/A 22-Sep-94 EA-2 54.08 N/A N/A 32.63 21.45 N/A 12-Oct-94 EA-2 54.08 N/A N/A 31.94 22.14 N/A 09-Sep-92 EA-3 54.32 N/A N/A 34.36 19.96 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A	21-Mar-94	EA-2	54.08					
02-Jun-94 EA-2 54.08 N/A N/A 36.86 17.22 N/A 07-Jun-94 EA-2 54.08 N/A N/A 36.70 17.38 N/A 28-Jun-94 EA-2 54.08 N/A N/A 36.18 17.90 N/A 07-Jul-94 EA-2 54.08 N/A N/A 34.71 19.37 N/A 11-Aug-94 EA-2 54.08 N/A N/A 31.42 22.66 N/A 22-Sep-94 EA-2 54.08 N/A N/A 32.63 21.45 N/A 12-Oct-94 EA-2 54.08 N/A N/A 31.94 22.14 N/A 01-Sep-92 EA-3 *N/A N/A N/A 34.36 19.96 N/A 09-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A	08-Apr-94	EA-2	54.08					
07-Jun-94 EA-2 54.08 N/A N/A 36.70 17.38 N/A 28-Jun-94 EA-2 54.08 N/A N/A 36.18 17.90 N/A 07-Jul-94 EA-2 54.08 N/A N/A 34.71 19.37 N/A 11-Aug-94 EA-2 54.08 N/A N/A 31.42 22.66 N/A 22-Sep-94 EA-2 54.08 N/A N/A 32.63 21.45 N/A 12-Oct-94 EA-2 54.08 N/A N/A 31.94 22.14 N/A 01-Sep-92 EA-3 *N/A N/A N/A 34.36 19.96 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A	19-May-94	EA-2	54.08					
28-Jun-94 EA-2 54.08 N/A N/A 36.18 17.90 N/A 07-Jul-94 EA-2 54.08 N/A N/A 34.71 19.37 N/A 11-Aug-94 EA-2 54.08 N/A N/A 31.42 22.66 N/A 22-Sep-94 EA-2 54.08 N/A N/A 32.63 21.45 N/A 12-Oct-94 EA-2 54.08 N/A N/A 31.94 22.14 N/A 09-Sep-92 EA-3 54.32 N/A N/A 34.36 19.96 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A N/A 34.39 19.93 N/A	02-Jun-94	EA-2	54.08					
07-Jul-94 EA-2 54.08 N/A N/A 34.71 19.37 N/A 11-Aug-94 EA-2 54.08 N/A N/A 31.42 22.66 N/A 22-Sep-94 EA-2 54.08 N/A N/A 32.63 21.45 N/A 12-Oct-94 EA-2 54.08 N/A N/A 31.94 22.14 N/A 01-Sep-92 EA-3 *N/A N/A N/A 34.60 N/A N/A 09-Sep-92 EA-3 54.32 N/A N/A 34.36 19.96 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A	07-Jun-94	EA-2						
11-Aug-94 EA-2 54.08 N/A N/A 31.42 22.66 N/A 22-Sep-94 EA-2 54.08 N/A N/A 32.63 21.45 N/A 12-Oct-94 EA-2 54.08 N/A N/A 31.94 22.14 N/A 01-Sep-92 EA-3 *N/A N/A N/A 34.60 N/A N/A 09-Sep-92 EA-3 54.32 N/A N/A 34.36 19.96 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A	28-Jun-94	EA-2	54.08					
22-Sep-94 EA-2 54.08 N/A N/A 32.63 21.45 N/A 12-Oct-94 EA-2 54.08 N/A N/A N/A 31.94 22.14 N/A 01-Sep-92 EA-3 54.32 N/A N/A N/A 34.60 N/A N/A 09-Sep-92 EA-3 54.32 N/A N/A 34.36 19.96 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A	07-Jul-94	EA-2						
12-Oct-94 EA-2 54.08 N/A N/A 31.94 22.14 N/A 01-Sep-92 EA-3 *N/A N/A N/A 34.60 N/A N/A 09-Sep-92 EA-3 54.32 N/A N/A 34.36 19.96 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A	11-Aug-94	EA-2						
01-Sep-92 EA-3 *N/A N/A N/A 34.60 N/A N/A 09-Sep-92 EA-3 54.32 N/A N/A 34.36 19.96 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A	22-Sep-94	EA-2						
09-Sep-92 EA-3 54.32 N/A N/A 34.36 19.96 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A		EA-2	54.08	N/A	N/A	31.94	22.14	N/A
09-Sep-92 EA-3 54.32 N/A N/A 34.36 19.96 N/A 13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A	04 8 02	E	*NI/A	N/A	N/A	34 60	N/A	N/A
13-Sep-92 EA-3 54.32 N/A N/A 34.39 19.93 N/A	•							
10 000 02	•							
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TABLE 4 SUMMARY OF GAUGING DATA FOR MILITARY GAS STATION ON MAIN BASE, EGLIN AFB, VALPARAISO, OKALOOSA COUNTY, FLORIDA.

	Weil	Casing	Depth	LPH	Depth To	Corrected	. LPH
Date	Number	Elevation	To LPH	Elevation	Water	Water Elev.	Thicknes:
		(feet MSL)	(feet)	(feet MSL)	(feet)	(feet MSL)	(feet)
21-May-93	EA-3	54.32	N/A	N/A	35.59	18.73	N/A
18-Jun-93	EA-3	54.32	N/A	N/A	36.48	17.84	N/A
27-Jul-93	EA-3	54.32	N/A	N/A	36.70	17.62	N/A
17-Aug-93	EA-3	54.32	N/A	N/A	36.73	17.59	N/A
16-Sep-93	EA-3	54.32	N/A	N/A	36.46	17.86	N/A
05-Oct-93	EA-3	54.32	N/A	N/A	36.78	17.54	N/A
11-Feb-94	EA-3	54.32	N/A	N/A	37.43	16.89	N/A
07-Mar-94	EA-3	54.32	N/A	N/A	36.35	17.97	N/A
21-Mar-94	EA-3	54.32	N/A	N/A	35.80	18.52	N/A
08-Apr-94	EA-3	54.32	N/A	N/A	35.99	18.33	N/A
19-May-94	EA-3	54.32	N/A	N/A	36.78	17.54	N/A
02-Jun-94	EA-3	54.32	N/A	N/A	37.11	17.21	N/A
07-Jun-94	EA-3	54.32	N/A	N/A	37.02	17.30	N/A
07-Jul-94	EA-3	54.32	N/A	N/A	35.36	18.96	N/A
11-Aug-94	EA-3	54.32	N/A	N/A	31.65	22.67	N/A
22-Sep-94	EA-3	54.32	N/A	N/A	32.84	21.48	N/A
12-Oct-94	EA-3	54.32	N/A	N/A	32.18	22.14	N/A
00 50= 02	E	55.47	N/A	N/A	34.97	20.50	N/A
09-Sep-92	EA-4	55.47 55.47	N/A	N/A	35.00	20.47	N/A
13-Sep-92	EA-4	55.47 55.47	N/A	N/A	35.22	20.25	N/A
22-Jan-93	EA-4	55.47 55.47	N/A	N/A	36.04	19.43	N/A
21-May-93	EA-4		N/A	N/A	36.93	18.54	
18-Jun-93	EA-4	55.47 55.47	N/A N/A	N/A	37.13		N/A
27-Jul-93	EA-4	55.47	N/A	N/A		18.34	N/A
17-Aug-93	EA-4	55.47	N/A N/A	N/A	37.25 37.00	18.22 18.47	N/A
16-Sep-93 05-Oct-93	EA-4	:55.47 55.47	N/A	N/A	37.33	18.14	N/A
11-Feb-94	EA-4	55.47 55.47	N/A	N/A	38.00	17.47	N/A N/A
07-Mar-94	EA-4	55.47 55.47	N/A	N/A	36.97	18.50	N/A
	EA-4		N/A	N/A	36.41		
21-Mar-94	EA-4	55.47 55.47	N/A	N/A	36.58	19.06 18.89	N/A N/A
08-Apr-94	EA-4 EA-4	55.47 55.47	N/A	N/A	37.31	18.16	N/A
19-May-94 02-Jun-94	EA-4	55.47 55.47	N/A	N/A	37.64	17.83	N/A
			N/A	N/A	37.55	17.92	N/A
08-Jun-94 29-Jun-94	EA-4	55.47 55.47	N/A	N/A	37.01	18.46	N/A
	EA-4	55.47	N/A	N/A	36.07	19.40	N/A
07-Jul-94	EA-4 EA-4	55.47	N/A	N/A	32.22	23.25	N/A
11-Aug-94		55.47	N/A	N/A	33.32	22.15	N/A
22-Sep-94 12-Oct-94	EA-4 EA-4	55.47 55.47	N/A	N/A	32.62	22.85	N/A
00 0 00	ea e	*NI/A	N/A	N/A	35.06	N/A	NI/A
02-Sep-92	EA-5	*N/A	N/A N/A				N/A
04-Sep-92	EA-5	*N/A		N/A	34.96	N/A	N/A
09-Sep-92	EA-5	55.24 55.24	N/A	N/A	34.88	20.36	N/A
13-Sep-92	EA-5	55.24 55.24	N/A	N/A	34.93	20.31	N/A
22-Jan-93	EA-5	55.24	N/A	N/A	35.17	20.07	N/A
21-May-93	EA-5	55.24	N/A	N/A	36.03	19.21	N/A
18-Jun-93	EA-5	55.24	N/A	N/A	36.95	18.29	N/A
27-Jul-93	EA-5	55.24	N/A	N/A	37.12	18.12	N/A
17-Aug-93	EA-5	55.24	N/A	N/A	37.16	18.08	N/A

TABLE 4 SUMMARY OF GAUGING DATA FOR MILITARY GAS STATION ON MAIN BASE, EGLIN AFB, VALPARAISO, OKALOOSA COUNTY, FLORIDA.

Date Number Elevation To LPH Elevation (feet) Water Elev Thickness (feet)	· 	Well	Casing	Depth	LPH	Depth To	Corrected	LPH
	Date							
16-Sep-93 EA-5 55.24 N/A N/A 38.91 18.33 N/A 05-Oct-93 EA-5 55.24 N/A N/A 37.25 17.99 N/A 11-Feb-94 EA-5 55.24 N/A N/A 37.91 17.33 N/A 07-Mar-94 EA-5 55.24 N/A N/A 37.91 17.33 N/A 07-Mar-94 EA-5 55.24 N/A N/A 36.75 18.49 N/A 21-Mar-94 EA-5 55.24 N/A N/A 36.75 18.49 N/A 08-Apr-94 EA-5 55.24 N/A N/A 36.31 18.93 N/A 08-Apr-94 EA-5 55.24 N/A N/A 36.44 18.80 N/A 08-Apr-94 EA-5 55.24 N/A N/A 36.44 18.80 N/A 02-Jun-94 EA-5 55.24 N/A N/A 37.57 17.67 N/A 08-Jun-94 EA-5 55.24 N/A N/A 37.57 17.67 N/A 08-Jun-94 EA-5 55.24 N/A N/A 37.57 17.67 N/A 29-Jun-94 EA-5 55.24 N/A N/A 37.45 17.79 N/A 29-Jun-94 EA-5 55.24 N/A N/A 35.86 19.38 N/A 07-Jun-94 EA-5 55.24 N/A N/A 35.86 19.38 N/A 11-Aug-94 EA-5 55.24 N/A N/A 35.86 19.38 N/A 12-Oct-94 EA-5 55.24 N/A N/A 33.30 21.94 N/A 12-Oct-94 EA-5 55.24 N/A N/A 33.30 21.94 N/A 12-Oct-94 EA-5 55.24 N/A N/A 35.47 20.38 N/A 12-Oct-94 EA-5 55.85 N/A N/A 35.47 20.38 N/A 12-Oct-94 EA-5 55.85 N/A N/A 35.47 20.38 N/A 12-Oct-94 EA-6 55.85 N/A N/A 35.47 20.38 N/A 13-Sep-92 EA-6 55.85 N/A N/A 35.47 20.38 N/A 13-Sep-92 EA-6 55.85 N/A N/A 35.47 20.38 N/A 13-N/A 37.49 EA-6 55.85 N/A N/A 37.40 18.40 N/A 37.40 N/A 37.40 N/A 37.40 N/A 37.40 N/	Date	110111201						
05-Oct-93 EA-5 55.24 N/A N/A 37.25 17.99 N/A 11-Feb-94 EA-5 55.24 N/A N/A 37.91 17.33 N/A 07-Mar-94 EA-5 55.24 N/A N/A 37.91 17.33 N/A 08-Apr-94 EA-5 55.24 N/A N/A 36.51 18.93 N/A 08-Apr-94 EA-5 55.24 N/A N/A 36.31 18.93 N/A 19-May-94 EA-5 55.24 N/A N/A 36.31 18.93 N/A 19-May-94 EA-5 55.24 N/A N/A 37.24 18.00 N/A 19-May-94 EA-5 55.24 N/A N/A 37.57 17.67 N/A 08-Jun-94 EA-5 55.24 N/A N/A 37.57 17.67 N/A 08-Jun-94 EA-5 55.24 N/A N/A 37.57 17.67 N/A 08-Jun-94 EA-5 55.24 N/A N/A 37.45 17.79 N/A 08-Jun-94 EA-5 55.24 N/A N/A 37.45 17.79 N/A 11-Aug-94 EA-5 55.24 N/A N/A 36.90 18.34 N/A 07-Jun-94 EA-5 55.24 N/A N/A 36.90 18.34 N/A 11-Aug-94 EA-5 55.24 N/A N/A 32.13 23.11 N/A 12-Oct-94 EA-5 55.24 N/A N/A 33.30 21.94 N/A 11-Aug-94 EA-5 55.24 N/A N/A 33.30 21.94 N/A 12-Oct-94 EA-5 55.24 N/A N/A 33.30 21.94 N/A 12-Oct-94 EA-5 55.24 N/A N/A 33.30 21.94 N/A 13-Sep-92 EA-6 55.85 N/A N/A 35.45 20.40 N/A 13-Sep-92 EA-6 55.85 N/A N/A 35.47 20.38 N/A 13-Jun-93 EA-6 55.85 N/A N/A 35.73 20.12 N/A 27-Jun-93 EA-6 55.85 N/A N/A 35.73 20.12 N/A 12-May-93 EA-6 55.85 N/A N/A 37.64 18.21 N/A 12-May-93 EA-6 55.85 N/A N/A 37.64 18.21 N/A 17-Aug-93 EA-6 55.85 N/A N/A 37.64 18.99 N/A 11-Feb-94 EA-6 55.85 N/A N/A 37.64 18.97 N/A 08-Apr-94 EA-6 55.85 N/A N/A 37.60 18.97 N/A 08-Apr-94 EA-6 55.85 N/A N/A 37.60 18.97 N/A 08-Apr-94 EA-6 55.85 N/A N/A 37.60 18.97 N/A 08-Apr-94 EA-6 55.85 N/A N/A 37.60 18.97 N/A 08-Apr-94 EA-6 55.85 N/A N/A 37.60 18.97 N/A 19-May-94 EA-6 55.85 N/A N/A 37.60 22.10-94 EA-6 55.85 N/A N/A 37			(,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			V-1-7		(.554)
05-Cet-93 EA-5 55.24 N/A N/A 37.25 17.99 N/A 11-Feb-94 EA-5 55.24 N/A N/A 37.91 17.33 N/A 07-Mar-94 EA-5 55.24 N/A N/A 38.791 17.33 N/A 07-Mar-94 EA-5 55.24 N/A N/A 36.71 18.93 N/A 08-Apr-94 EA-5 55.24 N/A N/A 36.31 18.93 N/A 19-May-94 EA-5 55.24 N/A N/A 36.31 18.93 N/A 19-May-94 EA-5 55.24 N/A N/A 37.24 18.00 N/A 19-May-94 EA-5 55.24 N/A N/A 37.57 17.57 N/A 08-Jun-94 EA-5 55.24 N/A N/A 37.57 17.57 N/A 08-Jun-94 EA-5 55.24 N/A N/A 37.57 17.57 N/A 08-Jun-94 EA-5 55.24 N/A N/A 37.45 17.79 N/A 19-May-94 EA-5 55.24 N/A N/A 37.45 17.79 N/A 19-May-94 EA-5 55.24 N/A N/A 37.45 17.79 N/A 11-Aug-94 EA-5 55.24 N/A N/A 36.90 18.34 N/A 11-Aug-94 EA-5 55.24 N/A N/A 33.30 21.94 N/A 11-Aug-94 EA-6 55.85 N/A N/A 33.30 21.94 N/A 11-Aug-94 EA-6 55.85 N/A N/A 33.30 21.94 N/A 11-Aug-94 EA-6 55.85 N/A N/A 35.47 20.38 N/A 11-Aug-94 EA-6 55.85 N/A N/A 35.47 20.38 N/A 11-Feb-94 EA-6 55.85 N/A N/A 36.54 19.31 N/A 12-1-May-93 EA-6 55.85 N/A N/A 36.54 19.31 N/A 11-Aug-93 EA-6 55.85 N/A N/A 37.64 18.43 N/A 11-Feb-94 EA-6 55.85 N/A N/A 37.64 18.21 N/A 11-Feb-94 EA-6 55.85 N/A N/A 37.60 18.97 N/A 08-Apr-94 EA-6 55.85 N/A N/A N/A 37.60 18.97 N/A 07-Apr-94 EA-6 55.85 N/A N/A N/A 37.60 18.97 N/A 07-Apr-94	16-Sep-93	EA-5	55.24	N/A	N/A	36.91	18.33	N/A
11-Feb-94 EA-5 55.24 N/A N/A 37.91 17.33 N/A 07-Mar-94 EA-5 55.24 N/A N/A 36.31 18.99 N/A 08-Apr-94 EA-5 55.24 N/A N/A 36.31 18.93 N/A 08-Apr-94 EA-5 55.24 N/A N/A 36.31 18.93 N/A 08-Apr-94 EA-5 55.24 N/A N/A 36.31 18.93 N/A 08-Apr-94 EA-5 55.24 N/A N/A 37.24 18.00 N/A 02-Jun-94 EA-5 55.24 N/A N/A 37.24 18.00 N/A 02-Jun-94 EA-5 55.24 N/A N/A 37.57 17.67 N/A 08-Jun-94 EA-5 55.24 N/A N/A 37.45 17.79 N/A 08-Jun-94 EA-5 55.24 N/A N/A 36.90 18.34 N/A 07-Jun-94 EA-5 55.24 N/A N/A 36.90 18.34 N/A 07-Jun-94 EA-5 55.24 N/A N/A 36.90 18.34 N/A 07-Jun-94 EA-5 55.24 N/A N/A 36.90 18.34 N/A 11-Aug-94 EA-5 55.24 N/A N/A 33.30 21.94 N/A 12-Oct-94 EA-5 55.24 N/A N/A 33.30 21.94 N/A 12-Oct-94 EA-5 55.24 N/A N/A 33.30 21.94 N/A 12-Oct-94 EA-5 55.85 N/A N/A 35.45 20.40 N/A 12-Sep-92 EA-6 55.85 N/A N/A 35.47 20.38 N/A 13-Sep-92 EA-6 55.85 N/A N/A 35.43 19.31 N/A 27-Jul-93 EA-6 55.85 N/A N/A 37.76 18.09 N/A 18-Jun-93 EA-6 55.85 N/A N/A 37.64 18.21 N/A 27-Jul-93 EA-6 55.85 N/A N/A 37.64 18.21 N/A 17-Aug-93 EA-6 55.85 N/A N/A 37.50 18.35 N/A 05-Oct-93 EA-6 55.85 N/A N/A 37.50 18.35 N/A 07-Mar-94 EA-6 55.85 N/A N/A 37.50 18.35 N/A 07-Mar-94 EA-6 55.85 N/A N/A 37.50 18.35 N/A 07-Mar-94 EA-6 55.85 N/A N/A 37.50 18.35 N/A 07-Mar-94 EA-6 55.85 N/A N/A 37.50 18.37 N/A 07-Mar-94 EA-6 55.85 N/A N/A 37.50 18.37 N/A 07-Mar-94 EA-6 55.85 N/A N/A 37.50 18.39 N/A 07-Mar-94 EA-6 55.85 N/A N/A 37.50 18.39 N/A 07-Mar-94 EA-6 55.85 N/A N/A 37.50 18.39 N/A 07-Mar-94 EA-6 55.85 N/A N/A 37.50 18.39 N/A 07-Mar-94 EA-6 55.85 N/A N/A 37.50 18.39 N/A 07-Mar-94 EA-6 55.85 N/A N/A 37.50 19.30 N/A 07-Mar-94 EA-6 55.85 N/A N/A 37.50 19.30 N/A 07-Mar-94 EA-6 55.85					N/A		17.99	
07-Mar-94 EA-5 55.24 N/A N/A 36.75 18.49 N/A 21-Mar-94 EA-5 55.24 N/A N/A 36.31 18.93 N/A 08-Apr-94 EA-5 55.24 N/A N/A 36.41 18.80 N/A 19-May-94 EA-5 55.24 N/A N/A 37.24 18.00 N/A 02-Jun-94 EA-5 55.24 N/A N/A 37.24 18.00 N/A 02-Jun-94 EA-5 55.24 N/A N/A 37.45 17.79 N/A 08-Jun-94 EA-5 55.24 N/A N/A 37.45 17.79 N/A 29-Jun-94 EA-5 55.24 N/A N/A 37.45 17.79 N/A 29-Jun-94 EA-5 55.24 N/A N/A 36.90 18.34 N/A 07-Jun-94 EA-5 55.24 N/A N/A 35.80 18.34 N/A 11-Aug-94 EA-5 55.24 N/A N/A 35.80 19.38 N/A 11-Aug-94 EA-5 55.24 N/A N/A 35.80 19.38 N/A 11-Aug-94 EA-5 55.24 N/A N/A 35.30 21.94 N/A 12-Oct-94 EA-5 55.24 N/A N/A 33.30 21.94 N/A 12-Oct-94 EA-5 55.24 N/A N/A 35.47 20.38 N/A 22-Jan-93 EA-6 55.85 N/A N/A 35.47 20.38 N/A 22-Jan-93 EA-6 55.85 N/A N/A 35.47 20.38 N/A 18-Jun-93 EA-6 55.85 N/A N/A 35.47 20.38 N/A 11-Aug-93 EA-6 55.85 N/A N/A 37.42 18.43 N/A 17-Aug-93 EA-6 55.85 N/A N/A 37.42 18.43 N/A 17-Aug-93 EA-6 55.85 N/A N/A 37.50 18.35 N/A 11-Feb-94 EA-6 55.85 N/A N/A 36.83 19.02 N/A 11-Feb-94 EA-6 55.85 N/A N/A 36.83 19.02 N/A 11-Feb-94 EA-6 55.85 N/A N/A 37.06 18.79 N/A 11-Aug-94 EA-6 55.85 N/A N/A 36.83 19.02 N/A 11-Feb-94 EA-6 55.85 N/A N/A 37.06 18.79 N/A 11-Aug-94 EA-6 55.85 N/A N/A 36.83 19.02 N/A 11-Feb-94 EA-6 55.85 N/A N/A 37.06 18.79 N/A 11-Aug-94 EA-6 55.85 N/A N/A 37.50 18.35 N/A 11-Aug-94 EA-6 55.85 N/A N/A 37.50 18.35 N/A 11-Aug-94 EA-6 55.85 N/A N/A 37.50 18.37 N/A 11-Aug-94 EA-6 55.85 N/A N/A 37.50 18.37 N/A 11-Aug-94 EA-6 55.85 N/A N/A 37.50 18.37 N/A 11-Aug-94 EA-6 55.85 N/A N/A 37.50 18.37 N/A 11-Aug-94 EA-6 55.85 N/A N/A 37.50 18.37 N/A 11-Aug-94 EA-6 55.85 N/A N/A 37.50 18.37 N/A 11-Aug-94 EA-6 55.85 N/A N/A 37.50 18.37 N/A 11-Aug-94 EA-6 55.85 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39 N/A N/A 37.50 18.39				N/A	N/A	37.91	17.33	
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19-May-94 EA-6 55.85 N/A N/A 37.81 18.04 N/A 02-Jun-94 EA-6 55.85 N/A N/A 38.13 17.72 N/A 08-Jun-94 EA-6 55.85 N/A N/A 38.06 17.79 N/A 29-Jun-94 EA-6 55.85 N/A N/A 37.52 18.33 N/A 07-Jul-94 EA-6 55.85 N/A N/A 36.67 19.18 N/A 08-Jul-94 EA-6 55.85 N/A N/A 36.67 19.18 N/A 11-Aug-94 EA-6 55.85 N/A N/A 36.21 19.64 N/A 11-Aug-94 EA-6 55.85 N/A N/A 32.69 23.16 N/A 22-Sep-94 EA-6 55.85 N/A N/A 33.82 22.03 N/A 12-Oct-94 EA-6 55.85 N/A N/A 33.12 22.73 N/A 09-Sep-92 EA-7 53.94 N/A N/A 34.20 19.74 N/A 13-Sep-92 EA-7 53.94 N/A N/A 34.62 19.32 N/A 22-Jan-93 EA-7 53.94 N/A N/A 34.62 19.32 N/A 21-May-93 EA-7 53.94 N/A N/A 35.52 18.42 N/A 18-Jun-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.59 17.35 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A N/A 36.65 N/A N/A 37.28 16.66 N/A N/A 11-Feb-94 EA-7 53.94 N/A N/A N/A 37.28 16.66 N/A								N/A
02_Jun-94 EA-6 55.85 N/A N/A 38.13 17.72 N/A 08_Jun-94 EA-6 55.85 N/A N/A 38.06 17.79 N/A 29_Jun-94 EA-6 55.85 N/A N/A 37.52 18.33 N/A 07_Jul-94 EA-6 55.85 N/A N/A 36.67 19.18 N/A 08_Jul-94 EA-6 55.85 N/A N/A 36.21 19.64 N/A 11_Aug-94 EA-6 55.85 N/A N/A 32.69 23.16 N/A 11_Aug-94 EA-6 55.85 N/A N/A 33.82 22.03 N/A 12_Oct-94 EA-6 55.85 N/A N/A 33.12 22.73 N/A 09_Sep-92 EA-7 53.94 N/A N/A 34.20 19.74 N/A 13_Sep-92 EA-7 53.94 N/A N/A 34.24 19.70 N/A 21_May-93	•							. N/A
08-Jun-94 EA-6 55.85 N/A N/A 38.06 17.79 N/A 29-Jun-94 EA-6 55.85 N/A N/A 37.52 18.33 N/A 07-Jul-94 EA-6 55.85 N/A N/A 36.67 19.18 N/A 08-Jul-94 EA-6 55.85 N/A N/A 36.67 19.18 N/A 11-Aug-94 EA-6 55.85 N/A N/A 32.69 23.16 N/A 11-Aug-94 EA-6 55.85 N/A N/A 32.69 23.16 N/A 22-Sep-94 EA-6 55.85 N/A N/A 33.82 22.03 N/A 12-Oct-94 EA-6 55.85 N/A N/A 33.12 22.73 N/A 13-Sep-92 EA-7 53.94 N/A N/A 34.20 19.74 N/A 13-Sep-92 EA-7 53.94 N/A N/A 34.24 19.70 N/A 22-Jan-93 EA-7 53.94 N/A N/A 34.62 19.32 N/A 21-May-93 EA-7 53.94 N/A N/A 34.62 19.32 N/A 18-Jun-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.59 17.35 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A 37.28 16.66 N/A								N/A
29-Jun-94 EA-6 55.85 N/A N/A 37.52 18.33 N/A 07-Jul-94 EA-6 55.85 N/A N/A 36.67 19.18 N/A 08-Jul-94 EA-6 55.85 N/A N/A 36.21 19.64 N/A 11-Aug-94 EA-6 55.85 N/A N/A 32.69 23.16 N/A 11-Aug-94 EA-6 55.85 N/A N/A 33.82 22.03 N/A 12-Oct-94 EA-6 55.85 N/A N/A 33.82 22.03 N/A 12-Oct-94 EA-6 55.85 N/A N/A 33.12 22.73 N/A 12-Oct-94 EA-7 53.94 N/A N/A 34.20 19.74 N/A 13-Sep-92 EA-7 53.94 N/A N/A 34.24 19.70 N/A 22-Jan-93 EA-7 53.94 N/A N/A 34.62 19.32 N/A 21-May-93 EA-7 53.94 N/A N/A 35.52 18.42 N/A 18-Jun-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 27-Jul-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.59 17.35 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A 37.28 16.66 N/A							17.79	N/A
07-Jul-94 EA-6 55.85 N/A N/A 36.67 19.18 N/A 08-Jul-94 EA-6 55.85 N/A N/A 36.21 19.64 N/A 11-Aug-94 EA-6 55.85 N/A N/A 32.69 23.16 N/A 12-Sep-94 EA-6 55.85 N/A N/A 33.82 22.03 N/A 12-Oct-94 EA-6 55.85 N/A N/A 33.12 22.73 N/A 09-Sep-92 EA-7 53.94 N/A N/A 34.20 19.74 N/A 13-Sep-92 EA-7 53.94 N/A N/A 34.24 19.70 N/A 22-Jan-93 EA-7 53.94 N/A N/A 34.62 19.32 N/A 21-May-93 EA-7 53.94 N/A N/A 35.52 18.42 N/A 18-Jun-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 27-Jul-93								N/A
08-Jul-94 EA-6 55.85 N/A N/A 36.21 19.64 N/A 11-Aug-94 EA-6 55.85 N/A N/A 32.69 23.16 N/A 22-Sep-94 EA-6 55.85 N/A N/A 33.82 22.03 N/A 12-Oct-94 EA-6 55.85 N/A N/A 33.12 22.73 N/A 12-Oct-94 EA-7 53.94 N/A N/A 34.20 19.74 N/A 13-Sep-92 EA-7 53.94 N/A N/A 34.24 19.70 N/A 22-Jan-93 EA-7 53.94 N/A N/A 34.62 19.32 N/A 21-May-93 EA-7 53.94 N/A N/A 35.52 18.42 N/A 18-Jun-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 27-Jul-93 EA-7 53.94 N/A N/A 36.59 17.35 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.59 17.35 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.33 17.61 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A 37.28 16.66 N/A								N/A
11-Aug-94 EA-6 55.85 N/A N/A 32.69 23.16 N/A 22-Sep-94 EA-6 55.85 N/A N/A 33.82 22.03 N/A 12-Oct-94 EA-6 55.85 N/A N/A 33.12 22.73 N/A 09-Sep-92 EA-7 53.94 N/A N/A 34.20 19.74 N/A 13-Sep-92 EA-7 53.94 N/A N/A 34.24 19.70 N/A 22-Jan-93 EA-7 53.94 N/A N/A 34.62 19.32 N/A 21-May-93 EA-7 53.94 N/A N/A 35.52 18.42 N/A 18-Jun-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 27-Jul-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.59 17.35 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A 37.28 16.66 N/A								N/A
22-Sep-94 EA-6 55.85 N/A N/A 33.82 22.03 N/A 12-Oct-94 EA-6 55.85 N/A N/A 33.12 22.73 N/A 09-Sep-92 EA-7 53.94 N/A N/A 34.20 19.74 N/A 13-Sep-92 EA-7 53.94 N/A N/A 34.24 19.70 N/A 22-Jan-93 EA-7 53.94 N/A N/A 34.62 19.32 N/A 21-May-93 EA-7 53.94 N/A N/A 35.52 18.42 N/A 18-Jun-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 27-Jul-93 EA-7 53.94 N/A N/A 36.59 17.35 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A N/A 36.65 17.29 N/A 11							23.16	N/A
12-Oct-94 EA-6 55.85 N/A N/A 33.12 22.73 N/A 09-Sep-92 EA-7 53.94 N/A N/A 34.20 19.74 N/A 13-Sep-92 EA-7 53.94 N/A N/A 34.24 19.70 N/A 22-Jan-93 EA-7 53.94 N/A N/A 34.62 19.32 N/A 21-May-93 EA-7 53.94 N/A N/A 35.52 18.42 N/A 18-Jun-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 27-Jul-93 EA-7 53.94 N/A N/A 36.59 17.35 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.33 17.61 N/A 05-Oct-93 EA-7 53.94 N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A 37.28 16.66 N/A	_						22.03	N/A
09-Sep-92 EA-7 53.94 N/A N/A 34.20 19.74 N/A 13-Sep-92 EA-7 53.94 N/A N/A 34.24 19.70 N/A 22-Jan-93 EA-7 53.94 N/A N/A 34.62 19.32 N/A 21-May-93 EA-7 53.94 N/A N/A 35.52 18.42 N/A 18-Jun-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 27-Jul-93 EA-7 53.94 N/A N/A N/A 36.59 17.35 N/A 17-Aug-93 EA-7 53.94 N/A N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A 37.28 16.66 N/A	•					33.12	22.73	N/A
13-Sep-92 EA-7 53.94 N/A N/A 34.24 19.70 N/A 22-Jan-93 EA-7 53.94 N/A N/A 34.62 19.32 N/A 21-May-93 EA-7 53.94 N/A N/A 35.52 18.42 N/A 18-Jun-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 27-Jul-93 EA-7 53.94 N/A N/A N/A 36.59 17.35 N/A 17-Aug-93 EA-7 53.94 N/A N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.33 17.61 N/A 05-Oct-93 EA-7 53.94 N/A N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A 37.28 16.66 N/A	12 000 04	2.0	•					•
13-Sep-92 EA-7 53.94 N/A N/A 34.24 19.70 N/A 22-Jan-93 EA-7 53.94 N/A N/A 34.62 19.32 N/A 21-May-93 EA-7 53.94 N/A N/A 35.52 18.42 N/A 18-Jun-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 27-Jul-93 EA-7 53.94 N/A N/A 36.59 17.35 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.33 17.61 N/A 05-Oct-93 EA-7 53.94 N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A 37.28 16.66 N/A	09-Sep-92	EA-7	53.94	N/A	N/A	34.20	19.74	N/A
22-Jan-93 EA-7 53.94 N/A N/A 34.62 19.32 N/A 21-May-93 EA-7 53.94 N/A N/A 35.52 18.42 N/A 18-Jun-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 27-Jul-93 EA-7 53.94 N/A N/A 36.59 17.35 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.33 17.61 N/A 05-Oct-93 EA-7 53.94 N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A 37.28 16.66 N/A				N/A	N/A	34.24	19.70	N/A
21-May-93 EA-7 53.94 N/A N/A 35.52 18.42 N/A 18-Jun-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 27-Jul-93 EA-7 53.94 N/A N/A 36.59 17.35 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.33 17.61 N/A 05-Oct-93 EA-7 53.94 N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A 37.28 16.66 N/A	•				N/A	34.62	19.32	N/A
18-Jun-93 EA-7 53.94 N/A N/A 36.39 17.55 N/A 27-Jul-93 EA-7 53.94 N/A N/A 36.59 17.35 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.33 17.61 N/A 05-Oct-93 EA-7 53.94 N/A N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A 37.28 16.66 N/A						35.52	18.42	N/A
27-Jul-93 EA-7 53.94 N/A N/A 36.59 17.35 N/A 17-Aug-93 EA-7 53.94 N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.33 17.61 N/A 05-Oct-93 EA-7 53.94 N/A N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A N/A 37.28 16.66 N/A	•		· ·				17.55	N/A
17-Aug-93 EA-7 53.94 N/A N/A 36.60 17.34 N/A 16-Sep-93 EA-7 53.94 N/A N/A 36.33 17.61 N/A 05-Oct-93 EA-7 53.94 N/A N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A 37.28 16.66 N/A						36.59	17.35	N/A
16-Sep-93 EA-7 53.94 N/A N/A 36.33 17.61 N/A 05-Oct-93 EA-7 53.94 N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A 37.28 16.66 N/A						36.60	17.34	N/A
05-Oct-93 EA-7 53.94 N/A N/A 36.65 17.29 N/A 11-Feb-94 EA-7 53.94 N/A N/A 37.28 16.66 N/A	-					36.33	17.61	N/A
11-Feb-94 EA-7 53.94 N/A N/A 37.28 16.66 N/A	•							N/A
1110001								
07-Mar-94 EA-7 53.94 N/A N/A 35.15 17.79 N/A	07-Mar-94	EA-7	53.94	N/A	N/A	36.15	17.79	N/A

TABLE 4 SUMMARY OF GAUGING DATA FOR MILITARY GAS STATION ON MAIN BASE, EGLIN AFB, VALPARAISO, OKALOOSA COUNTY, FLORIDA.

		Well	Casing	Depth	LPH	Depth To	Corrected	LPH
	ate	Number	Elevation	To LPH	Elevation	Water	Water Elev.	Thicknes
			(feet MSL)	(feet)	(feet MSL)	(feet)	(feet MSL)	(feet)
21-	Mar-94	EA-7	53.94	N/A	N/A	35.64	18.30	N/A
08-	-Apr-94	EA-7	53.94	N/A	N/A	35.85	18.09	N/A
	May-94	EA-7	53.94	N/A	N/A	36.66	17.28	N/A
	Jun-94	EA-7	53.94	N/A	N/A	36.98	16.96	N/A
	Jun-94	EA-7	53.94	N/A	N/A	36.90	17.04	N/A
	-Jul-94	EA-7	53.94	N/A	N/A	34.94	19.00	N/A
	Aug-94	EA-7	53.94	N/A	N/A	31.52	22.42	N/A
	Sep-94	EA-7	53.94	N/A	N/A	32.75	21.19	N/A
	Oct-94	EA-7	53.94	N/A	N/A	32.03	21.91	N/A
12	-000-34	LA-7	00 .34	IVA	1474	02.00	21.51	13/7
	Sep-92	EA-8	56.04	N/A	N/A	35.15	20.89	N/A
	Sep-92	EA-8	56.04	N/A	N/A	35.17	20.87	N/A
22-	Jan-93	EA-8	56.04	N/A	N/A	35.30	20.74	N/A
21-1	May-93	EA-8	56.04	N/A	N/A	36.09	19.95	N/A
18-	Jun-93	EA-8	56.04	N/A	N/A	36.95	19.09	N/A
27	-Jul-93	EA-8	56.04	N/A	N/A	37.14	18.90	N/A
17-	Aug-93	EA-8	56.04	N/A	N/A	37.34	18.70	N/A
	Sep-93	EA-8	56.04	N/A	N/A	37.13	18.91	N/A
05-	Oct-93	EA-8	56.04	N/A	N/A	37.46	18.58	N/A
	Feb-94	EA-8	56.04	N/A	N/A	38.13	17.91	N/A
	Mar-94	EA-8	56.04	N/A	N/A	37.11	18.93	N/A
	Mar-94	EA-8	56.04	N/A	N/A	36.56	19.48	N/A
	Apr-94	EA-8	56.04	N/A	N/A	36.74	19.30	N/A
	May-94	EA-8	56.04	N/A	N/A	37.41	18.63	N/A
	Jun-94	EA-8	56.04	N/A	N/A	37.76	18.28	N/A
	Jun-94	EA-8	56.04	N/A	N/A	37.70	18.34	N/A
	-Jul-94	EA-8	56.04	N/A	N/A	35.99	20.05	N/A
	Aug-94	EA-8	56.04	N/A	N/A	32.34	23.70	N/A
	Sep-94	EA-8	56.04	N/A	N/A	33.37	22.67	N/A
	Oct-94	EA-8	56.04	N/A	N/A	32.65	23.39	N/A
00	0 00	E4 0	EE 40	N/A	N/A	24.04	20.52	N/A
	Sep-92	EA-9	55.43			34.91	20.52	
	Sep-92	EA-9	55.43	N/A	N/A N/A	35.07	20.36	N/A N/A
	Jan-93	EA-9	55.43	N/A		34.63 35.39	20.80	
	May-93	EA-9	55.43 55.43	N/A	N/A	35.38	20.05	N/A
	Jun-93	EA-9	55.43	N/A	N/A	36.29	19.14	N/A N/A
	-Jul-93	EA-9	55.43 55.43	N/A	N/A	36.43 37.14	19.00	N/A N/A
	Aug-93	EA-9	55.43	N/A	N/A	37.14 37.00	18.29	N/A N/A
	Sep-93	EA-9	55.43	N/A	N/A		18.43	N/A N/A
	Oct-93	EA-9	55.43 55.43	N/A	N/A	37.34 37.95	18.09	N/A N/A
	Feb-94	EA-9	55.43	N/A	N/A		17.48 19.47	N/A N/A
	Mar-94	EA-9	55.43	N/A	N/A	36.96	18.47	
	Mar-94	EA-9	55.43	N/A	N/A	36.38	19.05	N/A
	Apr-94	EA-9	55.43	N/A	N/A	36.53 27.25	18.90	N/A
	May-94	EA-9	55.43	N/A	N/A	37.25	18.18	N/A
	Jun-94	EA-9	55.43	N/A	N/A	37.56	17.87	N/A
	Jun-94	EA-9	55.43	N/A	N/A	37.51	17.92	N/A
	Jun-94	EA-9	55.43	N/A	N/A	36.97	18.46	N/A
07	-Jul-94	EA-9	55.43	N/A	N/A	36.00	19.43	N/A

TABLE 4 SUMMARY OF GAUGING DATA FOR MILITARY GAS STATION ON MAIN BASE, EGLIN AFB, VALPARAISO, OKALOOSA COUNTY, FLORIDA.

·	Well	Casing	Depth	LPH	Depth To	Corrected	LPH
Date	Number	Elevation	To LPH	Elevation	Water	Water Elev.	Thickness
		(feet MSL)	(feet)	(feet MSL)	(feet)	(feet MSL)	(feet)
					(1.1.1)	(,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	(1000)
11-Aug-94	EA-9	55.43	N/A	N/A	32.20	23.23	N/A
22-Sep-94	EA-9	55.43	N/A	N/A	33.29	22.14	N/A
12-Oct-94	EA-9	55.43	N/A	N/A	32.63	22.80	N/A
18-Jun-93	EA-10	*N/A	N/A	N/A	36.63	N/A	N/A
27-Jul-93	EA-10	54.72	N/A	N/A	36.79	17.93	N/A
17-Aug-93	EA-10	54.72	N/A	N/A	36.87	17.85	N/A
16-Sep-93	EA-10	54.72	N/A	N/A	36.60	18.12	N/A
05-Oct-93	EA-10	54.72	N/A	N/A	36.91	17.81	N/A
11-Feb-94	EA-10	54.72	N/A	N/A	37.58	17.14	N/A
07-Mar-94	EA-10	54.72	N/A	N/A	36.49	18.23	N/A
21-Mar-94	EA-10	54.72	N/A	N/A	35.96	18.76	N/A
08-Apr-94	EA-10	54.72	N/A	N/A	36.15	18.57	N/A
19-May-94	EA-10	54.72	N/A	N/A	36.91	17.81	N/A
02-Jun-94	EA-10	54.72	N/A	N/A	37.26	17.46	N/A
07-Jun-94	EA-10	54.72	N/A	N/A	37.17	17.55	N/A
29-Jun-94	EA-10	54.72	N/A	N/A	36.60	18.12	N/A
07-Jul-94	EA-10	54.72	N/A	N/A	35.62	19.10	N/A
11-Aug-94	EA-10	54.72	N/A	N/A	31.81	22.91	N/A
22-Sep-94	EA-10	54.72	N/A	N/A	32.96	21.76	N/A
12-Oct-94	EA-10	54.72	N/A	N/A	32.29	22.43	N/A
18-Jun-93	EA-11	*N/A	N/A	N/A	35.90	N/A	N/A
27-Jul-93	EA-11	53.58	N/A	N/A	36.10	17.48	N/A
17-Aug-93	EA-11	53.58	N/A	N/A	36.07	17.51	N/A
16-Sep-93	EA-11	53.58	N/A	N/A	36.81	16.77	N/A
05-Oct-93	EA-11	53.58	N/A	N/A	36.13	17.45	N/A
11-Feb-94	EA-11	53.58	N/A	N/A	36.76	16.82	N/A
07-Mar-94	EA-11	53.58	N/A	N/A	35.61	17.97	N/A
21-Mar-94	EA-11	53.58	N/A	N/A	35.12	18.46	N/A
08-Apr-94	EA-11	53.58	N/A	N/A	35.33	18.25	N/A
19-May-94	EA-11	53.58	N/A	N/A	36.13	17.45	N/A
02-Jun-94	EA-11	53.58	N/A	N/A	36.44	17.14	N/A
07-Jun-94	EA-11	53.58	N/A	N/A	36.31	17.27	N/A
07-Jul-94	EA-11	53.58	N/A	N/A	34.29	19.29	N/A
11-Aug-94	EA-11	53.58	N/A	N/A	30.98	22.60	N/A
22-Sep-94	EA-11	53.58	N/A	N/A	32.25	21.33	N/A
12-Oct-94	EA-11	53.58	N/A	N/A	31.53	22.05	N/A
24 Man 24	EA 40	*N/A	N/A	N/A	35.46	N/A	N/A
21-Mar-94	EA-12	*N/A	N/A	N/A	35.48	N/A	N/A
23-Mar-94	EA-12	53.96	N/A	N/A	35.63	18.33	N/A
08-Apr-94	EA-12 EA-12	53.96	N/A	N/A	36.43	17.53	N/A
19-May-94		53.96	NA	N/A	36.76	17.20	N/A
02-Jun-94	EA-12	53.96	N/A	N/A	36.58	17.28	N/A
07-Jun-94	EA-12	53.96	N/A	N/A	34.55	19.41	N/A
07-Jul-94	EA-12	53.96	N/A	N/A	31.30	22.66	N/A
11-Aug-94	EA-12	53.96	N/A	N/A	32.52	21.44	N/A
22-Sep-94	EA-12	53.96	N/A	NA	31.82	22.14	N/A
12-Oct-94	EA-12	JJ.30	144	IVA	J 1.02	44.17	1 1// 1

TABLE 4 SUMMARY OF GAUGING DATA FOR MILITARY GAS STATION ON MAIN BASE, EGLIN AFB, VALPARAISO, OKALOOSA COUNTY, FLORIDA.

D-4-	Well	Casing	Depth	LPH	Depth To	Corrected	LPH
Date	Number	Elevation	To LPH	Elevation	Water	Water Elev.	Thickness
		(feet MSL)	(feet) .	(feet MSL)	(feet)	(feet MSL)	(feet)
14-Mar-94	EA 12	*N/A	N/A	N/A	36.97	N/A	NI/A
	EA-13	*N/A	N/A	N/A	36.90	N/A N/A	N/A
21-Mar-94	EA-13			N/A			N/A
23-Mar-94	EA-13	*N/A	N/A		36.94	N/A	N/A
08-Apr-94	EA-13	54.78	N/A	N/A	37.02	17.76	N/A
19-May-94	EA-13	54.78	N/A	N/A	37.83	16.95	N/A
02-Jun-94	EA-13	54.78	N/A	N/A	38.14	16.64	N/A
07-Jun-94	EA-13	54.78	N/A	N/A	38.05	16.73	N/A
07-Jul-94	EA-13	54.78	N/A	N/A	36.08	18.70	N/A
11-Aug-94	EA-13	54.78	N/A	N/A	32.74	22.04	N/A
22-Sep-94	EA-13	54.78	N/A	N/A	34.02	20.76	N/A
12-Oct-94	EA-13	54.78	N/A	N/A	33.32	21.46	N/A
14-Mar-94	EA-14	*N/A	N/A	N/A	35.21	N/A	N/A
21-Mar-94	EA-14	*N/A	N/A	N/A	35.23	N/A	N/A
23-Mar-94	EA-14	*N/A	N/A	N/A	35.26	N/A	N/A
08-Apr-94	EA-14	53.46	N/A	N/A	35.46	18.00	N/A
19-May-94	EA-14	53.46	N/A	N/A	36.27	17.19	N/A
02-Jun-94	EA-14	53.46	N/A	N/A	36.59	16.87	N/A
07-Jun-94	EA-14	53.46	N/A	N/A	36.31	17.15	N/A
07-Jul-94	EA-14	53.46	N/A	N/A	33.34	20.12	N/A
11-Aug-94	EA-14	53.46	N/A	N/A	31.13	22.33	N/A
22-Sep-94	EA-14	53.46	N/A	N/A	32.41	21.05	N/A
12-Oct-94	EA-14	53.46	N/A	N/A	31.70	21.76	N/A
14-Mar-94	EA-15	*N/A	N/A	N/A	35.23	N/A	N/A
21-Mar-94	EA-15	*N/A	N/A	N/A	35.25	N/A	N/A
23-Mar-94	EA-15	*N/A	N/A	N/A	35.25	N/A	N/A
08-Apr-94	EA-15	53.41	N/A	N/A	35.46	17.95	N/A
19-May-94	EA-15	53.41	N/A	N/A	36.27	17.14	N/A
02-Jun-94	EA-15	53.41	N/A	N/A	36.60	16.81	N/A
07-Jun-94	EA-15	53.41	N/A	N/A	36.31	17.10	N/A
07-Jul-94	EA-15	53.41	N/A	N/A	33.79	19.62	N/A
11-Aug-94	EA-15	53.41	N/A	N/A	31.15	22.26	N/A
22-Sep-94	EA-15	53.41	N/A	N/A	32.44	20.97	N/A
12-Oct-94	EA-15	53.41	N/A	N/A	31.72	21.69	N/A
21-May-93	SVE-1	DRY AT 32.07					
18-Jun-93	SVE-1	DRY AT 32.07					
27-Jul-93	SVE-1	DRY AT 32.07					
29-Jun-94	SVE-1	DRY AT 32.07					
21-May-93	SVE-2	DRY AT 31.82					
18-Jun-93	SVE-2	DRY AT 31.82					
27-Jul-93	SVE-2	DRY AT 31.82					
27-Jun-93 29-Jun-94	SVE-2	DRY AT 31.82				•	
23-Jul 1-34	345-2	DIXI AI 51.02					
21-May-93	AAS-1	*N/A	N/A	N/A	35.94	N/A	N/A
18-Jun-93	AAS-1	*N/A	N/A	N/A	36.82	N/A	N/A

TABLE 4 SUMMARY OF GAUGING DATA FOR MILITARY GAS STATION ON MAIN BASE, EGLIN AFB, VALPARAISO, OKALOOSA COUNTY, FLORIDA.

Date	Well Number	Casing Elevation (feet MSL)	Depth To LPH (feet)	LPH Elevation (feet MSL)	Depth To Water (feet)	Corrected Water Elev. (feet MSL)	LPH Thickness (feet)
24 Jun 02	AAS-1	*N/A	N/A	N/A	36.95	N/A	N/A
21-Jun-93	AAS-1	55.21	N/A	N/A	37.00	N/A	N/A
27-Jul-93			· N/A	N/A	37.10	N/A	N/A
17-Aug-93	AAS-1	55.21				N/A	N/A
16-Sep-93	AAS-1	55.21	N/A	N/A	36.85		
05-Oct-93	AAS-1	55.21	N/A	N/A	37.21	N/A	N/A
07-Mar-94	AAS-1	55.21	N/A	N/A	36.78	N/A	N/A
21-Mar-94	AAS-1	55.21	N/A	N/A	36.24	N/A	N/A
08-Apr-94	AAS-1	55.21	N/A	N/A	36.44	N/A	N/A
19-May-94	AAS-1	55.21	N/A	N/A	37.17	N/A	N/A
02-Jun-94	AAS-1	55.21	N/A	N/A	37.51	N/A	N/A
08-Jun-94	AAS-1	55.21	N/A	N/A	37.40	N/A	N/A
29-Jun-94	AAS-1	55.21	N/A	N/A	36.86	N/A	N/A
07-Jul-94	AAS-1	55.21	N/A	N/A	35.91	N/A	N/A
11-Aug-94	AAS-1	55.21	N/A	N/A	32.06	N/A	N/A
22-Sep-94	AAS-1	55.21	N/A	N/A	33.20	N/A	N/A
12-Oct-94	AAS-1	55.21	N/A	N/A	32.51	N/A	N/A

^{*}N/A = Elevation of index on top of casing had not been established as of this date.

TABLE 5 ORGANIC VAPOR ANALYZER (OVA) HEADSPACE ANALYSIS OF SOIL SAMPLES COLLECTED AT THE MILITARY GAS STATION ON THE MAIN BASE, EGLIN AFB, VALPARAISO, OKALOOSA COUNTY, FLORIDA.

Drilling Location	Depth (feet)	Total Hydrocarbon Headspace Reading (ppm)	Filtered Headspace (Methane only) (ppm)	Corrected Headspace ⁽⁴⁾ (ppm)
				(ррш)
EA-1	5-7	2.6		0.0
	7-9	2.6	· —	0.0
	9-11	2.6		0.0
	11-13	2.6	. _	0.0
	13-15	2.8	-	0.2
	15-17	4.5	3.2	1.3*
	17-19	4.0	3.4	0.6
	19-21	4.0	3.2	0.8
	21-23	4.0	4.0	0.0
	23-25	3.4	3.0	0.4
	25-27	5.2	4.2	1.0
	27-29	4.0	3.5	0.5
	29-31	4.1	· 3.5	0.6
	31-33	3.8	3.2	0.6
	33-35	2.8	_	0.2
	35-37	3.0	_	0.4
	40-42	3.6	3.0	0.6
EA-2	5-7	1.8	_	0.0
	10-12	2.3	-	0.3
	15-17	1.8		0.0
	20-22	2.8	2.4	0.4
	25-27	3.5	2.2	1.3*
	30-32	2.0		0.2
	35-37	4.4	3.2	1.2
	40-42	3.2	2.8	0.4
EA-3	5-7	5.6	_	0.3
	7-9	5.6	coulom	0.1
•	9-11	5.8	5.6	0.3
	11-13	5.8	5.6	0.2
	: 13-15	6.0	5.7	0.3
	15-17	5.8	5.6	0.2
	17-19	6.6	6.0	0.6
	19-21	5.8	-	0.2
	21-23	6.0		0.4
	23-25	5.6	_	0.0
	25-27	5.8		0.2
	27-29	6.8	. 6.2	0.6
	29-31	6.7	6.3	0.4
	31-33	6.0	5.8	0.2
	33-35	7.0	6.2	0.8
	40-42	7.2	6.3	0.9

[©] Corrected OVA headspace determined by subtracting the ambient background reading (not shown) and the filtered (methane-only) reading from the total hydrocarbon headspace reading.

from the total hydrocarbon headspace reading.

— = Methane concentration not evaluated due to low initial headspace reading.

NR = No sample recovery.

* Indicates sample sent to lab for analysis.

[&]quot;Soil sample collected from cuttings.

TABLE 5 (Cont.)

Drilling Location	Depth (feet)	Total Hydrocarbon Headspace Reading (ppm)	Filtered Headspace (Methane only) (ppm)	Corrected Headspace ^(A) (ppm)
EA-4	5-7	6.0		0.4
1	10-12	5.5	-	0.0
1	15-17	6.6	6.2	0.4
1	17-19	7.0	6.1	0.9
II .	19-21	6.6	6.0	0.6
1	21-23	22.0	8.6	
li .	23-25	7.1	6.2	13.4
1	25-27	110.0		0.9
l			22.0	0.88
	27-29	220.0	55.0	165.0
{ }	29-31	1000+	550.0	450+
	31-33	1000+	480.0	520+°
	33-35	1000+	620.0	380+
l	35-37	1000+	510.0	490+
J	40-42	850	340.0	510.0
			3-10.0	210.0
EA-S	5-7	2.4		0.0
	10-12	2.6		0.0
l	15-17	2.6		
l .				0.2
	20-22	3.0	2.6	0.4
1	25-27	2.5	_	0.1
	30-32	3.2	2.7	0.5*
	35-37	5.9	2.9	3.0
	40-42	3.0	2.7	0.3
	. ~			
EA-6	5-7	1.5	-	0.0
	10-12	1.5	_	0.0
1	15-17	1.4	_	0.0
	20-22	2.4	1.8	0.6*
•	25-27	1.7		0.2
	30-32	2.1	1.7	0.4
	35-37	1.6	_	0.1
	40-42	2.8	1.8	1.0
EA-7	5-7	3.3		0.0
	10-12	3.3	_	0.0
	15-17	3.4	_	0.1
	20-22	3.9	3.5	0.4
	25-27	3.6	_	0.1
	30-32	3.8	_	0.2
	35-37	3.6	_	0.2
	40-42	3.8	. 3.6	. 0.2
EA-8	5-7	4.2	_	0.0
MA W.	10-12	4.2		0.0
	15-17	4.2	_	0.0
	20-22	4.2	_	0.0
	25-27	4.2	-	0.0
	30-32	4.2	_	0.0*
	35-37	4.2		0.0
	40-42	4.4		0.2
				li li
EA-9	43-45	22.0	5.6	16.4
	53-55	30.0	14.0	29.0
EA-10	. 4-6	1.5	_	0.0
	9-11	2.0	1.5	0.5
	14-16	5.2	1.6	3.6
	19-21	2.8	1.6	1.2
	24-26	3.6	1.6	2.0
	29-31		2.0	
		6.8	2.0	4.8
	34-36	21.0	2.0	19.0°
	39-41	NR	NR	NR
	44-46	76.0	15.0	59.0

TABLE 5 (Cont.)

Drilling Location	Depth (feet)	Total Hydrocarbon Headspace Reading (ppm)	Filtered Headspace (Methane only) (ppm)	Corrected Headspace ^(s) (ppm)
EA-11	4-6	7.0	6.9	0.1
	9-11	6.8	6.6	0.2
	14-16	6.8	6.7	0.1
	19-21	6.8	6.7	0.1
	24-26	7.3	7.0	0.3
	29-31	7.2	6.9	0.3*
	34-36	8.4	7.3	1.1
	39-41	210.0	70.0	133.0
	44-46	NR	NR	NR
EA-12	1-1.25	5.3	_	0.0
	2-2.4	5.6		0.0
	3-3.5	5.8		0.0
	4-4.1	5.8	_	0.0
	5-7	5.8	-	0.2
	7-9	6.0		0.0
	9-11	5.9	·	0.0
	15-17	6.0	· ·	0.0
	20-22	6.0		0.0
	25-27	6.0	_	0.0
	30-32	6.2	_	0.0
	35-37	6.2	_	0.0
	40-42	6.2		0.0
EA-13	1-1.1	0.6		0.0
	2-2.2	0.6		0.0
	3-3.5	0.6		0.0
	4-4.1	0.6	-	0.0
	5-7	0.6 0.6	-	. 0.0
	9-11 15-17	0.6	_	0.0 0.0
	20-22	0.6	_	0.0
	25-27	0.6		0.0
	30-32	0.6	_	0.0
	35-37	0.6	denice	0.0
	40-42	0.6	_	0.0
EA-14	1-1.4	0.3		0.0
	2-2.8	0.4		0.0
	3-3.4	0.4	_	0.0
	4-4.3	0.4		0.0
	5-7	0.4		0.0
	7-9	0.4	-	0.0
	9-11	0.4	_	0.0
	15-17	0.4	-	0.0
	20-22	0.4	-	0.0
	30-32	0.4	_	0.0
	35-37 40-42	0.4 0.4	Ξ	0.0 0.0
			_	
EA-15	1-1.2	0.3	-	0.0
	2-2.3	0.3		0.0
	3-3.1	0.2 0.2	_	0.0 0.0
	4-4.3 5-7	0.3		0.0
	3-7 7-9	0.3	_	0.0
	9-11	0.3	_	0.0
	15-17	0.3	-	0.0
	20-22	0.4	-	0.0
	25-27	0.3	_	0.0
	30-32	0.3	_	0.0
	35-37	0.3	_	0.0
	40-42	0.3		0.0
	50 ^{ee}	0.3		0.0 -
	59**	0.3		0.0

TABLE 5 (Cont.)

Drilling Location	Depth (feet)	Total Hydrocarbon Headspace Reading (ppm)	Filtered Headspace (Methane only) (ppm)	Corrected Headspace ⁽⁴⁾ (ppm)
A	4-6	0.7	_	0.0
	6-8	1.3	1.1	0.2
	8-10	0.8		0.1
	10-12	0.9	_	0.2
1	15-17	1.3	1.0	0.3
	20-22	1.1		0.2
ł	25-27	1.2	1.0	0.2
	30-32	1.0		0.1
	35-37	1.4	1.4	0.0
	40-42	110.0	10.0	100.0
В	4-6	0.4	_	0.2
	6-8	0.2	_	0.0
f	8-10	0.3	_	0.2
	10-12	0.6		0.4
	15-17	0.2	_	0.0
	20-22	70.0	25.0	45.0
	25-27	210.0	117.0	93.0 •
l .	27-29	50.0	5.2	44.8
	29-31	56.0	3.0	53.0
	31-33	10.0	5.5	4.5
	33-35	12.0	5.6	6.4
	36-38	700.0	38.0	662.0
	40-42	72.0	9.4	62.6
С	5- 7	3.8	_	0.2
	7-9	3.8		0.0
•	9-11	4.0		0.2
	15-17	4.1	_	0.1
	20-22	4.4	4.2	0.2
	25-27	5.2	4.8	0.4
	30-32	4.6	4.3	0.3
	35-37	4.6	4.2	0.4
	40-42	430.0	14.0	416.0
D	: 5-7	3.8	-	0.2
	10-12	3.8	_	0.0
	15-17	3.8		0.0
	20-22	4.0	-	0.2
	25-27	4.3	. 4.0	0.3
	30-32	3.6	3.4	0.2
	35-37	3.2	<u></u>	0.4
	40-42	3.8	3.4	1.0
E	5-7	1.3		0.0
	10-12	1.5		0.2
	15-17	4.4	2.2	2.2
	20-22	2.4	2.0	0.4
	25-27	2.9	2.2	0.7
	30-32	2.6	2.0	0.6
	35-37	3.1	2.5	0.6
	40-42	2.2	1.8	0.4

TABLE 5 (Cont.)

Drilling Location	Depth (fæt)	Total Hydrocarbon Headspace Reading (ppm)	Filtered Headspace (Methane only) (ppm)	Corrected Headspace ^ω (ppm)
7	5-7	5.6	_	0.0
1	7-9	6.0	5.8	0.2
	9-11	6.1	5.8	0.3
		5.6	3.8	0.0
	11-13	6.0	5.7	0.0
•	13-15	6.1		
	15-17	5.8	5.7	0.4
	17-19			0.2
	19-21	7.2	6.2	1.0
	21-23	6.7	6.1	. 0.6
	23-25	5.9	5.7	0.2
	25-27	6.0	5.8	0.2
	27-29	8.1	6.4	1.7
	29-31	6.9	6.2	0.7
l	31-33	25.0	10.0	15.0
l l	33-35	5.9	_	0.3
	40-42	15.0	8.6	. 6.4
ĸ	5-7	4.2	-	0.4
	7-9 ·	4.1	-	0.3
	9-11	4.8	4.2	0.6
	11-13	62.0	12.0	50.0
	13-15	19.0	8.5	10.5
	15-17	110.0	22.0	88.0*
	17-19	28.0	10.0	18.0
	19-21	100.0	23.0	77.0
	21-23	14.0	5.5	8.5
	23-25	95.0	18.0	77.0*
	25-27	77.0	16.0	61.0
•	27-29	32.0	11.0	21.0
	29-31	34.0	13.0	21.0
	31-33	47.0	9.5	37.5
	33-35	20.0	7.8	12.2
	35-37	15.0	5.0	10.0
	40-42	12.0	6.2	5.8
L	5-7	3.0	_	0.0
	10-12	3.4	-	0.2
	15-17	3.6	3.2	0.4
	20-22	3.3	08940	0.3
	25-27	4.0	3.3	0.7
l	30-32	4.4	· 3.4	1.0
l	35-37	6.4	3.8	2.6
	40-42	3.8	3.4	0.4
Q	4-6	2.6	· _	0.0
ı `	6-8	2.6	_	0.0
	8-10	2.7	2.6	0.1
	10-12	2.6		0.0
	12-14	2.7	2.6	0.1
i	14-16	3.0	2.8	0.2*
1	16-18	2.9	2,8	0.1
	18-20	2.9	2.9	0.0
	20-22	3.1	3.0	0.1
1	24-26	3.0	-	0.0
	29-31	3.0		0.0
	34-36	3.2	3.1	0.1

TABLE 5 (Cont.)

	Depth (feet)	Total Hydrocarbon Headspace Reading (ppm)	Filtered Headspace (Methane only) (ppm)	Corrected Headspace ⁽¹⁾ (ppm)
			NA	0.0
R	4-6	6.6	NA ·	0.0
	6-8	7.0 13.0	6.8 7.4	0.2 5.6
	8-10	7.3	7.0	0.3
	10-12	7.3 15.0	7.0 7.2	7.8
	12-14 14-16	7.4	7.0	0.4
	16-18	13.0	7.6	5.4
	18-20	14.0	8.0	6.0
	24-26	74.0	27.0	47.0*
	29-31	8.4	7.4	1.0
	34-36	14.0	8.2	5.8
		5.4	_	0.0
S	4-6 6-8	5.4	_	0.0
	8-10	5.5	5.4	0.1
	10-12	5.8	5.5	0.3
	12-14	5.6	5.5	0.1
	14-16	6.4	5.7	0.7
	16-18	6.9	5.8	1.1
	18-20	6.9	5.8	1.1*
	24-26	5.9	5.6	0.3
	29-31	6.2	5.6	0.6
	34-36	5.9	5.6	0.3
т	4-6	4.4		0.0
•	6-8	4.5	4.4	0.1
	8-10	4.6	4.4	0.2
	10-12	4.4		0.0
i e	12-14	4.4		0.0
	14-16	4.5	_	0.1
	16-18	4.5	4.4	0.1
	18-20	4.6	4.4	0.2
	24-26	5.4	4.6	0.8
	29-31	6.8	5.5	1.3*
	34-36	10.0	5.4	4.6
ט	4-6	6.0	_	0.0
	6-8	6.0	_	0.0
	8-10	5.7		0.0
	10-12	6.2	5.8	0.4
ł	12-14	5.8	· 5.7	0.1
	14-16	5.9	5.7	0.2
	16-18	6.0	5.8	0.2
	18-20	6.2	5.8	0.4 1.5
	20-22	7.4 7.4	5.9 5.8	1.6
	22-24	6.0	5.8 5.7	0.3
	24-26 26-28	6.2	5.8	0.4
1	28-30	9.4	5.8	3.6
l	28-30 30-32	15.0	6.0	9.0
	30-32 32-34	9.2	6.2	3.0
	34-36	68.0	12.0	56.0

TABLE 5 (Cont.)

Drilling Location	Depth (feet)	Total Hydrocarbon Headspace Reading (ppm)	Filtered Headspace (Methane only) (ppm)	Corrected Headspace ⁽⁴⁾ (ppm)
v	4-6	5.0		0.0
Y	6-8	6.0		0.0
	8-10	7.1	_	0.0
	10-12	7.4	_	0.0
	12-14	100.0	10.0	90.0
	14-16	90.0	9.0	81.0
1	16-18	220.0	18.0	202.0
	18-20	120.0	15.0	105.0
1	20-22	590.0	22.0	568.0
1	22-24	1000+	32.0	1000+7
	24-26	1000+	40.0	1000+
	26-28	1000+	45.0	1000+
	28-30	1000+	52.0	1000+
1	30-32	940.0	30.0	910.0
1	32-34	1000+	45.0	1000+*
	34-36	960	26.0	934.0
W	4-6	7.2	_	0.0
	6-8	7.3	7.2	0.1
	8-10	7.4	7.2	0.2
i	10-12	7.5	7.3	0.3
1	12-14	7.2		0.0
	14-16	7.2	-	0.0
	16-18	7.6	7.2	0.4*
	18-20	7.2		0.0
	24-26	7.6	7.3	0.3
	29-31	7.4	7.2	0.2
	34-36	7.4	7.2	0.2
SVE-1	24-26	1.2	(0)	0.4 ^(c)
SVE-2	24-26	3.8	3.8	O(e)
AAS-1	4-6	0.6	0.6	0
LVV7.1	9-11	0.8	0.2	0.4
	14-16	0.2	0.2	0
	19-21	0.2	0.2	0
	24-26	0.8	0.2	0.4
	29-31	1.2	0.2	0.6
	34-36	2.6	0.2	1.4
	39-41	600	37	565 ^(*)
	44-46	380	4.0	372
	49-51	780	2.8	772
	54-56	120	0	115
	59-61	16	0	11

(ABLE 6: Summary of Analytical Data Results for Soil Samples Collected at the Military Gas Station Eglin AFB, Vaparaiso, Okaloosa County, Florida.

	Parameter		TRPH	ТРН	Total Kjeldahl Nitrogen	Total Nitrate	Total Ortho- Phosphate	Ammonia Nitorgen	Total Hydrocarbon Degraders
	Method		9073	418.1	356.2	300	300	350.1	9215B
	Unit		mg/Kg	mg/L	mg N/Kg	mg/Kg	mg/Kg	mg N/Kg	Colonies/ML
***	Detection limit		5	1	12.4-12.6	2.1	2.0-2.2	2.6-2.9	1
Date	Samipe I.D.	Depth				Concentrati	ion		
8/19/92	Ea-1	15-17	BDL .						
through	Ea-2	25-27	BOL						
9/1/92	EA-3	27-29	BOL						
	EA-4	31-33	BDL						
	EA-5	30-32	BDL						
	EA-6	20-22	BDL						
	EA-7	20-22	BDL						
	EA-8	30-32	BDL						
	K	15-17	BOL						
	EA-1(Dup)	15-17	BOL						
į.	Equipment Blank	-		BOL					
									:
*				•					
11/5/93	SVE-1	24-26		40.2	20	7.9	BDL	6.89	BDL
through	SVE-2	24-26		BDL	11.4	6.2	BDL	BOL	BOL
5/13/93	AAS-1	39-41		40.8	BDL	6.9	BDL	BOL	11
	AAs-1	49-51		-	BDL	BOL	BDL	BOL	10
	`.								
				·					
6/14/93	EA-10	34-36		BOL	•		•		
through	EA-11	29-31		BOL					
17/6/93	Q	14-16		BOL					
	R	24-26		BOL					
	s	18-20		BDL					
	T	29-31		BOL					
	U	30-32		BDL					
	V	10-12		BDL					
	· ·			BOL					
	V	32-34							
-	v w	16-18		BOL					
	V	1							

BDL Dup Below Detection Limit Sample duplicate

.-S.J.AMARY OF ANALYTICAL RESULTS FOR GROUNDWATE. SAMPLES COLLECTED AT THE MILITARY GAS ! ON THE MAIN BASE, IRP SITE ST-58, EGLIN AFB, VALPARAISO, OKALOOSA COUNTY, FLOIUDA. TABLE 4

					I.W.	WELL IDENTIFICATION AND SAMPLE DATE	ATION AND	SAMPLE DATI	
	EPA		Florida Target				EA-1 13	قدر المرابية فامنان	نا ب
Analyte	Method Nuniber	MDL"	Cleanip Concentration**	Units	9/13/92	66/91/6	+ 161718	11/26/96	74/01/2
Yelstilk Oreanic Aramatka"									
Benzene	602/8020	0.6.1	-	78,4	9'0>	4.1	⊽	⊽	⊽
Tohese	602/4010	_	RA	7/34	₹	-	1.1	V	⊽
Edrylbenzase	607/4010	1-60	HA	7,14	<0.9	07	9	⊽	⊽
Xylence, Total	602/8020	1-60	Y X	7,14	<0.0	543.6	320	~	ī
Total VOA	607/1020	1.9.0	20	, LT.	90>	696.5	481.7	~	Ţ
Medryl Test-Busyl Ester	209	5-25	2	724	₽	<15	!	Ţ	~
1,3. Dkhlorobeczene"	602/4020	1-1.1	RA	41/1	<.i.	<u>-</u>	•	-	⊽
1.4-Dichlombenzenen	602/4020	_	42	#E/L	~	~	2.9	V	V
1,2-Dichlerofenunce	601/1620	_	42	77.	ī	~		7	ī
Chlarobenizae	602/18020	1.1.3	٧×	1,31	<1.3	-	⊽	7	·
Palraudiese Aronaule Urdencerbene			;	:	,	;	,		
Naphibalene	0179/00/19/019	2.	*	7/17	•	<u>~</u>		~	₹
1-Methylnsphilistere	610/8100/820	07-1	٧×	#\$/L	į	ı	•	ī	₹
2-Methylaspirihaleac	610/8100/8110	01:1	٧×	121	<1.2	11	:	1.2	1.7
Toui Nipinhikae	610/8100/8270	1-10	8	real	<1.2	78	8.1	1 2	2.2
Accasphibyicae	610/8100/8270	01-1	NOC	784	c 3.1	2 2	-	₹	-
Acetaphilhene	610/8100/8270	0.3-10	MOL	7,31	'	2 2	_	⊽	5.3
Puercise	610/8100/8270	0.1-10	NDL	7/3/	<0.3	9	~	~	-
Phenanthrene	610/8100/8270	0.2-10	MDL	JISH.	<0.7	₽,	-	<u>-</u>	-
2,4-Dimethylphenel	0100/0019	2	XX	ug/L	į	Q >	:	:	!
Dibenzie, it jantlieze ene	610/1100/1270	0 4-10	MOL	re/L	*0 *	9 V	ī	⊽	Ţ
Anthracene	610/8100/8270	0.2-10	MDL	17A	< 0.2	= >	⊽	7	₹
Volatife Organic Halocarbane									
Terrelibracione	AUFO	-	NA NA	. 1/34	~	:	:	:	:
1, 4-Dichlurabenzene	0107	-	Y.	7/8/	<u>-</u>	:	:	:	:
1,1.Dichlarabentenc	0101	-	N.A.	ug/L	⊽	:	1	:	:
1.3.Dichlarobentene	0100	-	* *	7/3#	7	!	i	:	!
1,1-Dibremocthane ED3	204	0.03	0.02	יער	<0.0>	1	:	1	!
1,2.Dichloraethane	0108/109	-	2	הניר	⊽	i	ŧ		:
Total Petroleum Hydrocarban	-11	-	NA TPPIII 5 MIS	mg/L	V	:	:	:	:
				•					
Lead .	239.2	s	S. C. S.	# ! /L	₽	!	i	:	;
			· ·		,				

MDL = Mathed Detection Limit. The MOLs are given as a range for some analytes due to differences in dilution and methodologies over the time interval represented by the data.
 The Florida Target Chasup Concentrations were entrated from Section 17-770-730(\$Xa)(2), F.A.C.
 Only those VOAs detected in any sample are shown.
 A physical properties former concentrations are suspect as Method 40.20 does not employ a ballogen detector in confirm the presence of habitetized compounds.
 Only those VOH's detected in any sample are shown.
 Only those VOH's detected in any sample are shown.

^{*} The laboratory reported a negative bias for this sample. Actual concentrations may have been higher than thuse reported.

NA * No applicable Plurida target eleanup enneentrathur.

TABLE 4 (~ JIII.)

					WELL	WELL IDENTIFICATION AND SAMPLE DATE	THE SAMPL	E DATE
	EPA		Morida Target			w	EA-2	
Analyte	Medied	MDL	Chanup Concentration*1	Units	9/13/92	86/91/6	617194	6/11/94
Vakille Orzenic Aromaiket								
Benzene	001/8/109	0.6.1	-	חפינר	<0.6	~	7	~
Taluene	602/8070	_	< z	יוניר	⊽	<u>v</u> (-	⊽	~
Dhyhantene	001/1010	1.6.0	XX	7	<0.0>	017	₹	-
Xykaca, Total	607/1030	0.9.1	٧2	1121	<0.9	1.180	₹	7
Total VOA	007/1030	0.6.1	S	1/ 3 H	9.0 ×	1,520	₹	, n
Methyl Tert-Bulyl Elice	703	5-25	\$0	μΕ/I.	\$	<25	: ;	:
1.3 Dichlerobenicae"	601/8010	-:-	₹ 2	red.	<1.1	1>	Ţ	~
I. 4. Dichilerobenizate.	602/2030	-	4 2	78,	•	~	Ţ	7
1.2. Dichlerobentene	001/109	_	44	7/34	<u>-</u>	₹	-	V
Chlorobenzene	0101709	F1.5	٧,	7/80	<.i.	-	₹	~
Pohrwdrae Aromaik Higtocarbant"								
Naphhalene	07.18/10/18/10/19	1.10	Y Z	FE'L	<1.4 <1.4	÷	⊽	~
I-Mahylmphihalenc	610/8100/8270	0 <u>-</u> 1	ζ,	1,8/L	;	1	:	~
2. Madry losphilistene	610/8100/3270	0:1	< 2	7,5%	<1.7	2	:	:
Total Naphikalene	610/8100/12/0	DI-1	5	7,21	<1.7	93	~	~
Aconsplishingtone	610/1/00/17/0	0:-	MDL	787	<u>:</u>	01 >	~	⊽
Accesplishene	0111/0011/019	0.3-10	MDL	7/34	<1.7	= >	Ş	1.5
Puorene	610/8100/8270	0.2.10	NOL	787	<03	2 >	Ţ	~
Plensintere	61w11mu1220	0 2.10	MOL	1/30	< 0.2	01 ×	~	~
2.4. Directly plicing	810018	S	**	HE/L	:	< 10 < 10	:	:
Dilew(1,h) tohracete	610/8100/8270	0.4.10	MDL	7/3r	<0.4	× 10	~	J
Anthreene	610/B100/8170	0.2-10	MIDL	7/34	<0.1	01 >	7	~
Volatific Oceanic Habermont"								
Torreliorections	0101	-	*	784	-	:	:	i
1.4-Dichlare trans	DIOI	-	Y.	אניר	~	:	:	:
1,1.Dichturahourene	DIN	-	٧,	Jr3H	~	:	:	:
1.3-Dichlosbenzene	0102	-	Y.	1,8"	<u>~</u>	. :	:	:
1,2-Vilbrametihane	\$04	0.03	0 02	HVL.	< 0.02	:	:	፥
1,2 Okchierocihane	0108/109	-	1	אפור	⊽	:	:	:
Total Petraleum Aydrocarban	411.1		۲ ۲	T/Jus	₹	!	ı	ı
per			;	,	•			

TABLE 4 (wint.)

					WELL IDEN	WELL IDENTIFICATION AND SAMPLE DATE	WPLE DATE
	EPA		Flarida Target	-		EA-J	
Analyte	Method	MDL"	Cleniup Concentration"	Units	9/13/92	6/191/6	16/1/34
Seigible Organic Argmatica	603/1030	99.90	-	1/3r	9'0>	₹	₹
Tolliens	0008709	81-	N.	7/8/1	7	~	₹
Filmiltenzene	602/4010	8.6.0	Y.	MIN	<0.9	• =	₹
Yulence Test	602/4030	0.6.0	×2	ינור	<0.0	~	₹
Topi VOA	602/1020	0.6.60	20	7,31	9.0>	~	<u>~</u>
McAvi Ten-Buvi Elber	503	5.300	20	T/SH	≎	<u>.</u>	i
1.3. Dichloshenzene	602/1030	1-110	Y.V	F. J. C.	61.1	₹	₹
1.4.Dichlorohe mornel	602/1010	1-160	**	1/8×	⊽	<u>~</u>	⊽
1.2-Dichlosparene	602/8020	1-330	**	7/3/	₹	<u>~</u>	<u>-</u>
Chlorobensens	602/4020	1-130	¥	MEA	<1.3	₹	₹
Bellement to a second to the s							
Name of the state	610/1100/1270	92-1	YX	1/A/r	<1.4	<u>0</u> 1 >	~
1-Meibylosophibalene	610/8100/8770	27	. 42	7,37	i	:	•
2-Methylasobshatene	610/8100/8270	1-10	* Z	7,91	<1.1	01×	!
Taul Nichhikas	610/8100/8770	01-1	001	784	<1.2	oi >	⊽
Accorditylene	610/8100/1270	07-1	MDL	11 T	<3.1	o! >	-
Acenshorae	610/8100/4170	0.3-10	MDL	1,8/L	<1.1	01 >	1.9
Fluorane	610/8100/4270	0.3.10	MDL .	7/11	<0.3	01>	7.4
Phenatheen	610/8100/1270	0.2.10	MDL	71,311	<0.7	01>	<u>.</u>
2.4. Dimeraviohenoi	\$100/8270	0 2-10	4 Z	7,81	!	0; •	1
Discarda Mandricene	0128100/8270	0.4-10	MDL	M1/L	₹0>	0 >	⊽
Anthracene	0/18/00/8/019	0.2.10	MDL	7,811	<0.2	01>	⊽
Volette Organic Halocarbonia							
Tetraciloraechane	601/1010	-	≺ Z		~	i	!
1.4.Dichiatabeanes	0101/109	-	< Z	71,34	₹	:	!
1,2. Dichtorobenzene	0101/109	-	¥	ng/L	7	:	i
1,3-Othromeethane	8	0.02	0.02	"EL	< 0.03	:	:
			•	•	•		
1,1-Dichleroethane	01/1010	1-100	•	7,21	v	ì	:
Total Petroleam Mydrocarbon	18.1	-	NA	mg/L	7	i	:
Lead	239.2	~	51	11.21	\$:	ı

TABLE 4 (wint.)

						3	LL IDENTIF	ICATION AN	WELL IDENTIFICATION AND SAMPLE DATE	2	
	EPA		Florida Target					EA.4	·		
Analyte	Method	MDL	Cleanth Concentration ⁴⁴	Units	9/14/92	6,191/6	8/8/94	(1211)	1/20/96	11/1/196	2/10/97
Yolatile Organic Arematika**											
Benzone	601/8010	0.6-60	-	1181	3 4 ×	3.5	V	v	~	V	~
Talvene	602/8020	001-1	¥	7/8/	01	=	12	: _	. ₹	; ₹	₹
Chylbenzeuc	601/8010	0.9.90	¥	41/	25	\$10	*	310	4.00	2	740
Xyknct, Tetal	602/8010	0.6.0	4 2	מניר	4.900	2,340	2,100	1820	200	1010	1590
Tetal VOA	07.017.09	0.6-60	2	7/37	5,950	2,904.5	2,563	21.31	1540	9	2
Mellyl Tat-Buryl Edice	602	\$.50n	3.	Ž	< 500 000 000 000 000 000 000 000 000 000	× ×	:	V	V	~	J
1,3-Dichierobenzeechi	601/1010	011-	*	7/3/	VII0	270	ī,	v	7	V	~
I.4-Dichlerobenzenen	0001709	<u>3</u>	¥Z	7/8/	< 100)	31.	œ	•	•	-	~
1,3-Dichlorobenitoe	601/8020	1-330	¥	7/ 3 //	001 ×	< 330	120	ī	⊽	~	v
Chlorobenzene	601/8010	1:130	Y X	7/811	<130 <130	<u>~</u>	6.1	-	⊽	⊽	⊽
Polyneclese Acemails livdinessibans"											
Nepinfielais	610/1100/12/n	1.20	ž	75.	061	5	2	300	0.7	=	77
I-Maliylmplitlalane	610/11/00/12/20	1.20	YZ Z	7/211	:	:	!	1901	=	2.9	49
2-Maly insplutistone	610/8100/8370	<u>-</u>	Ϋ́	1/34	9	130	!	20	*	X.5	R
Total Naphthelene	610/1100/1270	01:1	20	7/30) (300	75	6.10	303	22.4	258
Accrepitity lene	610/1100/1270	9:	MDL	7/24	*	01.	:	58	2.1	-	5 ~
Acenaphiliene	410/8/00/8270	0.3.10	MDL	11811	<1.7	= >	э	25	~	3.2	3.5
Fluorenc	610/8100/8270	0.3-10	MDL	7/34	1.7	a. >	(.5	46		5.0	7.1
Phena athrene	610/8100/8270	0.2-10	NDL	7/30	3.5	01 >	4.2	=	.0%	1.2	~
7.4 Dimethylphonol	81004270	0.2-10	٧,	191	:	91 ×	:	:	:	:	:
Dibens(e,h)anthracene	410/8100/8270	0.4-10	NDL	17.	<0.4	01 >	~	01 >	v	7	ī
Anthraceae	610/11/00/11/10	0.7-10	MDL	ř	<02	01>	7	o 10	<u>,</u>	₹	₹
Yolaille Orsank Halacarlants											
Tetrachlomediane	. 0101/109	_	¥	787	<u>~</u>	:	:	:	:	:	:
1,4.Dichlorabeinene	6/1/101/1	-	YY X	7,27	-	:	i	:	i	;	:
1.3.Dichlotobeireue	01/109	-	××	7,3,1	ī	፡	i	:	:	:	:
1,3-Dibromechane	26	0.03	0.03	P\$/L	<0.02	!	•	:	:	:	i
1, 2-Dichbrucchane	0108/109	001:1	•	מוןר	00 ×	!	:	:	:	:	i
	•	•	;								
Tetal Petroleum Hydrocarbon	418.1	-	X X	nı V.L	=	ł	:	:	:	:	i
Lead	239.2	•	2	1,1,1	=	i	!	:	3	:	i
				•							

TABLE 4 (Lont.,

		· ·			WELL IDENTIF	WELL IDENTIFICATION AND SAMPLE DATE	PLE DATE
	EPA	-	Fluids Targo			EA-5	
Analyte	Number	NDL:	Cheanup	Unis	(4/1/6	16/91/6	16/11/9
Yelstille Orrenic Acomalice"							
Benuat	601/1010	0.6-1	-	ייפיר	9.0>	⊽	7
Tolura	003/109	-	4 Z	7,81	⊽	⊽	⊽
Ethy (beatene	602/18/120	1.60	KN KN	, E /I.	<0 >	⊽	⊽
Xylenes, Total	601/8010	1.6.0	YX	7/3rd	6.0>	₹	·
Total VOA	602/2020	1.9.0	50	. 1/3n	9.11>	⊽	⊽
Methyl Tert-Buyl Edier	2139	. ~	25	124	≎	\$ >	į
1,3-Dichlorobenzence	003/8030		42	7097	<1.1>	~	J
1,4.Dichlorebeneench	0001709	-	YX.	7/3/1	⊽	⊽	₹
1,2. Diehlorabenzenet	602/8020	_	42	7/31	⊽	₹	7
Chlorebeneeme	01011109	1.1.3	٧,	14.	<1.3	<u>-</u>	<u>.</u>
Polynucical Assaulte Hydrogathous!							
Naphhaleac	619/8106/8270	01-1	ž	uell	F1>	610	~
1-Melly Inshibalene	610/4100/4270	=======================================	~ 7	1/40	: :	i	; ;
2 - Methy lozabilisticae	610/8100/8220	- 10	7	1/00	<13	01>	
Tout Naphthakne	610/8100/8170	2:1	9		~ 7 >	01>	· ·
Acenaphihylene	610/8100/8270	9.1	M	med.		97	7
Acensphiese	610/8100/1170	0.3.10	NDI		C 1 2	97.7	; •
Fluorence	610/8100/1270	1-10	TOW TOWN	The state of		2 2	; 7
Phenauthreme	610/3100/1270	0.2.10	TON:	nv(l.	<0.7	01>	7
2,4-Dintechylphenul	1100/1270	0.2-10	ž	7	!	01>	:
Dibent(e,li)sadiracene	610/4100/8270	0.4-10	MDL	7/34	×0×	* I >	7
Amhracane	0128/0018/019	0 2-10	MDL	181r	< 0.2	× 10	~
Yellik Organ Halgertwart							
Terachine	0108/109	_	42	W/L	. ▼	:	:
1,4-Dichlorobentene	01011/109	_	٧z	77.	~	***	•
1,2.Dichlorobenzene	0101/100	-	Y.V	,.E/L	₹	!	!
1,2-Dibromecibane	304	0.03	0 03	7/3"	<0.02	•	i
1,2-Dkblaracibene	010\$/109	-	•	#1/L	<u>-</u>	į	;
Total Petraleum Hydrocarbon	418.1	-	NA	Ing/L	⊽	i	i
Lead	219.2	5	22	Was.	V	:	:
	!		?	1	;		

					WELL IDEN	WELL IDENTIFICATION AND SAMPLE DATE	AMPLE DATE
	EPA		Florida Target			EA-6	
Analyte	Number	MDL₩	Cleanup Concentration®	Units	8/14/92	66/91/6	16/11/9
Yelstik Ortenis Aramalist"							
Benunt	601/1030	1.9.0	-	MUL	9.0>	~	V
Toluene	0401/109	-	٧٧	we'll	₹⊽	; 7	; 7
Ethylbenzene	0207/109	0.9.1	H.	-	60>	7 7	7 7
Xylenes, Total	0104/209	14.0	Y Z	-	60>	; 7	; 7
Toul VOA	601/19020	0.6-1	2	-Jan	402	; 7	7 7
Methyl Teri-Butyl Ether	602	•	os	7,10	; V	: 7	;
1,3-Dichlorobenzene	0702/209	===	Y.V	#UL	; ;	; 7	: 7
L. C. Dichlorolentere	602/4010	-	*	7	~	; 🔻	7
1,1-Dichlotobenzeae	0201/109	-	ž	1/37	: 7	; 7	; ;
Chlorebengene	602/16010	14.3	¥	7	;;; ;;;	; ⊽	; ⊽
Polyauchar Aromatic Hydrocarbons							
Nephbukme	610/8100/8270	4	*2	<i>y</i>		41,	,
1-Methylosoluhalese	610/8100/11/20		2 2		¥. 1. V	200	<u>-</u>
2. Methylasphip bee	OCT BOOLENIA		۲:	7/17	•	!	:
Total Name of the	01100011017	2	Y .	7/84	<1.7	9 V	:
Accessioning	0.7800181017	2:	8	16	<1.1	01 >	⊽
Acertain	0/15/0015/015	0:-	HOL	1/A	<.1.L	01 >	,
	0/79/0019/019	0.1-10	Mol	7/31	<1.7	01 ×	₹
	010/8100/1/0	01-1	MDL	7,3,4	· ('0')	9 V	~
3 4 Direction Case	0/0/0/19/0/19	0.7.60	MDL	7,11	< 0.2	9	~
A. C. Curaciny anenal	1100/6270	0.2.10	¥	7,34	i	0:>	:
Ulbertele, A) souh sethe	610/8100/8179	0.4.10	אפר	7/3/1	+ 0>	< 10	~
Andriecine	610/1100/110	0.2.10	MDL	7/3"	<01	< 10	<u>~</u>
Yalatile Organic Halpenthere							
Teuschloradune	0108/109	_	**	J.***	7		
1,4-Dichlerobenzene	0101/109	_	* * Z		; ;	: :	i
1,2.Dichierobenases	010#/109	_	YX	Ž	₹∇	! !	i
I,1-Dibronochane	204	0 03	0.02	יהו	. 20:0>	ı	i
1.3 Deblarathan	0108/109	-	•		;		
		-	•	ייער	₹	!	i
Total Petroleum Hydrocarban	418.1	-	NA	n.g./L	₹	i	:
Lead	219.2	~	57	ne/L	V	i	;
		•	1	<u>.</u>	?	i	:

TABLE 4 Cont.,

Parish Early Parish Early Concention Parish Parish Concention Parish							WELL ID	ENTIFICAT	TION AND S	WELL IDENTIFICATION AND SAMPLE DATE	
Modeled MoDele Concentration Units 91(492)		EFA		Florida Targer					EA.7 12 12 44	1 1 1 1 m	
Colinois	Analyte	Mohud Nuaher	₩DL**	Cleanup Concentration**	Usitis	9/14/92	8/16/93	, 191713	A . 170176	11/15/96	79WIL
Column	Yoliith Orank Ammilia								•		
MA	Benune	602/8020	1.90	_	1/31	4.0>	Ţ	·	7	-	7
COLINGTO COLINGTO	Tolucia	601/8020		4 Z	7,81	-	⊽	ī	~	7	⊽
CONTROLOGY CON	Ellylbentese	601/8020	0 9.1	< %	7,31	6 N >	-	•	ī	~	~
60218020 0 6-1 51 60 60 60 60 60 60 60 60 60 60 60 60 60	Xylenes, Total	601/11/10	0.9.1	4 2	11211.	6.U.	⊽	7.4	⊽	-	-
602 1 1 1 1 1 1 1 1 1 4 1 1 1 1 1 1 1 1 1	Total VOA	602/8020	1-9:0	\$	7/50	9.0>	<u>~</u>	16.4	~	-	,
SOLIND	Melbyl Tert-Butyl Ether	209	. ~	20	12,	Ş	⊽	!	<u>~</u>	⊽	V
SOLISO10	1.3-Dicilorobenceisca	602/6020	==	42	72.	-1.1	V	7	⊽	7	<u>-</u>
SOZIONO	1.4-Dichlosebenzene*	602/8020	_	¥2	JIII.	<u>~</u>	<u>~</u>	1.1	~	<u>-</u>	-
610/80100 1-13 NA	1,2-Dichlesebenzere**	602/4010	-	¥×	PE/L	-	-	13	<u>,</u>	⊽.	~
STATE COLUMN STATE STA	Chlorobenzene	007/1030	1:1	٧X	rp/L	<1.1	₹	⊽	~	V	V
September 1-10	Polynucies Acomote Redressibant										
Sequence 1.10	Naphtialene	610:8100/8370	9:	¥×	1,2,1	4.1A	2	1.5	⊽	23	₹
Signation/1700 1.3-140 NA CG/L CG/L	1-Mcdiyasphilatene	610/81(x)/870		42	7/31	:	:	:	:	4.5	7.7
1-10 100	2-Methylnaphidialne	610/4100/470	1.1.10	4 2	7,5,1	< 1.2	= >	* * * * * * * * * * * * * * * * * * * *	·	£.5	₹
SOURTION SOURT S	Tout Naphthalene.	610/1100/13/70	9:-	100	75,	<1.1 <1.1	= >	 	<u>~</u>	~	2.3
1-10 MOL PELL CO.3	Accasplulytene	610/8100/820	<u>•</u>	MDL	701	C).	9 : V	<u>6.</u> ,	⊽:	·	<u>.</u>
100/12/12 100	Accuspinhenc	640/1100/1270	0 9	HOL	7.57	2.5	2 2	- :	₹ ₹	V 7	- 7
10 10 10 10 10 10 10 10	Fluorenc	0178/0018/019	91:0	102	724	()	2 5	? ?	7.	7 7	, •
Fig. 10 Fig.	7 4 Discharge	00000000	N. 7.10	701		, !		7 :	; ;	; ;	;
610/3100/1270 610/3100/1270 610/3100/3270 1-10 601/8010 1	Caroline and property of the control	or the least	2 7 6			707	2 5	V	V	V	7
601/R010 1 HA	Application	610/2100/220	0.2.10	MDF	1	<0.7	01 >	₹	V	V	⊽
601/R010 1 NA 11E/L <1 (211/R010 1 NA 11E/L <1 (211/R010 1 NA 11E/L <1 504 0.02 0.02 11E/L <0.02 601/8010 1 3 12E/L <1	Fluoranthene	610/8100/8270	1-10	MDL	7,34	⊽	01 ×	7	⊽	₹	9.
601/8010 1 HA 112/L <1 601/8010 1 HA 112/L <1 504 0 02 0.02 112/L <0.02 601/8010 1 3 112/L <1	Voletile Organic Habstarbonz"										
(418.1 1 NA 1.5/1. <1 column	Tetrachlomediane	601/8010	-	4 2	7,3,1	⊽	:	!	:	:	:
504 0 0? 0.02 1.g/L <1 climate	1.4-Dichluroleu/one	(201780)	_	44	1, g.1.	•	:	:	:	:	:
504 0 0 0 2 0.02 1/2 < 0.02 601/6010 1 3 1/2 < 1 < 1 < 1 ** ** ** ** ** ** ** ** ** ** ** ** *	1,2.Dkinforubenzene	0108/109	_	¥	<i>''</i>	⊽	:	;	:	:	:
601/8010 I 3 µg/L <1	1,2 Obromadhane	504	0 03	0.02	1/ 2 //	<0.02	i	i	:	:	:
418.1 1 HA nig/L <1	1,1.DicMarocibine	0109/109	_	3	7811	·	i	ŧ	1	:	i
	Total Petroleum Hydrocarbon	418.1	-	¥	ng/L	•	i	:	i	:	:
Lead 339.2 \$ 15 µz/L <\$	Lead	239.2	•	15	1/2×	~	i	!	:	i	፧

TABLE 4 (Lun.

						a dire moderation	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
	4				WELL IDEN	WELL IDENTIFICATION AND SANFLE DATE	ANIFLE DATE
	Medua		Cleanup			3	
Analyte	Number	MDL"	Concentration**	Units	9/14/92	9/16/93	P6/#/9
Yaluille Organic Argustica"							
Bentene	007/709	1.9.0	-	MOL	9.0>	~	17
Tolvane	0104/109	-	*	7 m	~	. ₹	7
Ethy lb era enc	00787100	14.0	< z	ne/L	60>	~	₹
Xyknes, Total	602/8020	0.9.1	٧٧	1/2"	60>	7	7
Total VOA	602/8020	1.9.0	20	1/30	<0.0>	₹	; -
Medinyl Test- Busyl Edher	un en	~	20	אפער	\$	*	; :
1,3-Diditorobenzene"	601/8010		4%	אניר.		. ≏	V
I.4. Dichlorobentene	601/8010	-	٧×	7/2/	-	~	Ţ
1,7.Dichlosobenitage	601/8020	_	₹2	7/84	<u>~</u>	₹	~
Chlaichentene	020\$/209	==	* 2	7/4"	<1.3	7	•
Petraucker Arematic Medicurathoni"							
Naphhakne	610/11/00/1270	9:-	< 2	ugil.	* ! >	=>	~
1- Methylaspialisteue	610/11/00/12/10	1.10	KN	7/3/	:	:	:
1. Nichylnspinstere	610/1100/1270	1.2.10	42	HE/L	<1.7	01 ×	:
Total Nephrinicae	610/8100/8270	<u> </u>	93	₩./L	~ >	UI >	~
Accordington	01078/0018/019	2	MDC	MB/L	<3.1	01 >	⊽
Acchapanen	610/6100/6270	2	MPL	7/3"	<1.1	= >	~
רוצחופשט	610/4100/8270	0:-	MDL	7/3"	<0.1	oi >	Ţ
	610/1100/1270	0.7.10	MDL	.1/34	<0.7	e >	7
I. 4- Dienethy phonoi	£100/£170	2	< z	יניר	:	01 ×	:
Dabene(a.h)saibtacene	010/8/00/8/019	0 + 0	MOL	1/1r	<0.4	o1 >	<u>~</u>
Anihitean	610/8100/8/20	0.2-10	MDL	7,17	< 0.2	212	⊽
Yolalik Orzanic Halocarbons							
Tetrachierachane	0101/109	_	4 2	7/8/	~	:	:
I,4-Dicition-benzeite	60174010	_	4 2	1/3/1	₹	:	:
I,7-Dichlerobentene	DIOK/IN9	_	4 X	ייניר	₹.	:	:
1,2.Dibremecthane	504	0.02	0.03	721	<0.0>	ŀ	;
1,2.Dichlorecthrac	0109/109	.		J. J.	⊽	i	i
Total Petroleum Hydrocarban	418.1	-	NA	T) Su	₹	ŧ	!
Lead	239.2	~	22		Š	i	:
				•			

TABLE 4 (~ont.,

					WELL IDEN	WELL IDENTIFICATION AND SAMPLE DATE	MPLE DATE
	EPA		Florida Target			EA:9	
Analyte	Medied Number	MDL"	Concentration*	Units	9/14/192	(8/9)/6	6/8/9/
Velatific Qreenis Aremetica"	603/2/200	0.6-1	_	ue/t.	9.0>	⊽	1
Telian	600/7003	-	ž	יייי	~	. △	~
Fibelbenzene	602/1030	1.6.0	YX	Mg/L	<0.9	⊽	v
Xylenes, Total	602/4010	0.9.7	XX	הוור	2.4	₹	·
Total VOA	601/16010	1-9:0	20	11 TH	7.4	~	⊽
Methyl Terl-Bulyl Ether	603	.1.25	\$	FB1F	≎	⊽,	!
1,3.Dkhlorobeareacta	601/1010	1-1-1	4 Z	PEAL.	l.1>	.⊽	₹
1,4-Dichlare benzene	603/1010		YZ:	7,8rd	7	₹ ;	· ;
1,3-Dichlers beazene" Chloroberraere	602/8020	- 1-1	* * *	֓֞֞֝֞֞֞֞֞֞֞֞֞֞֞֞֞֞֞֝֞֞֞֞֞֞֞֞֞֞֞֝֞֞֝֞֝֞֞֞֝	\ !!>	⊽ ⊽	7 7
				•			
Palynecker Aramaik Medrecarbon	616316A181613	91.1	42	li e :	7	9	ī
Maphibitas			< ₹ 2		; :	?, i	; !
2. Methy leaphthalene	610/100/12/10	2 : 1	: Z	7,47	~!>	9	:
i - Methylanehalene	610/8100/8270	01-1	Y X	17/1	!	:	:
Total Nephalene	610/8100/8270	01-1	901	1/1/n	<1.7	< 10	⊽
Acanaphylene	610/6100/8270	1.10	MDC	1/1×	<3.1	<10	<u>~</u>
Accomplishene	610/8100/1270	1.10	MDL	7,8/L	· 1.1>	<10	~
Fluorane	610/8100/8270	01:1	MDL	7,81	<0.3	01>	-
Phenaditene	610/1100/1270	1.10	NDL	7/811	<0.1	01>	-
2,4-Dimethylphenol	B100/8270	2	MA	7/3/	:	01×	:
Dibent(1,h)anthracene	610/8100/8270	0-1	MDL	"VL	×0.4	01>	V :
Anhitectic	610/8100/8170	0:1	MOL	ž	<0.7	₽:	-
Hearthbrobutsdiene	0121/00/11	2	₹	7/21	!	=	:
Yolathe Organic Helocardson							
Tetrachloroethene	0104	-	Y.Y	7/811	-	:	i
1,4-Dichlorobenzene	0100	-	4 X	7,80	V	i	:
1,2.Dichlerobeacene	0101	_	4	1/3×	-	:	:
1.3.Dichlorabentene	010		≺ Z	#1/F	:	•	:
1,2 Othermed bane	304	0.03	ω ο	, et	<0.02	:	:
1,2.Dichlaroethane	01011/109	2.	•	7/3"	~	!	:
Total Peroleum Hydrocarben	418.1	0.6-1	¥X	mg/L	-	!	:
Lead	239.2	23	15.	#1/F	\$	i	:

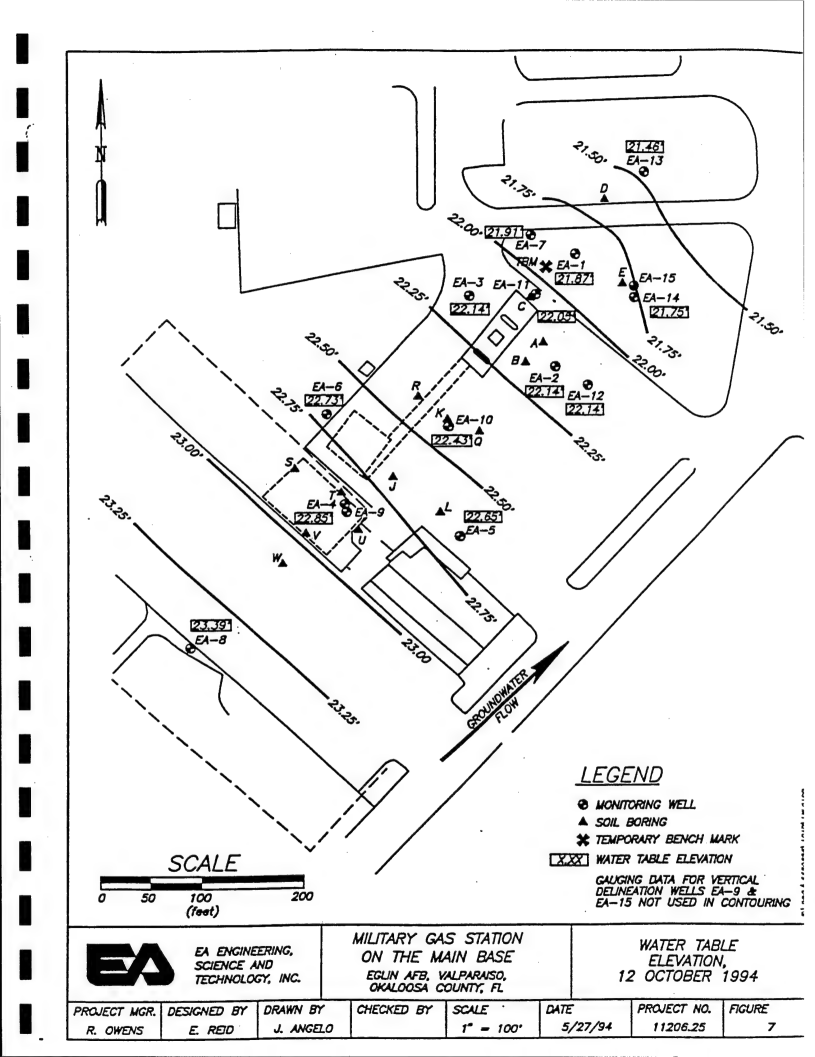
						WELL	WELL IDENTIFICATION AND SAMPLE DATE	ON AND SAR	IFLE DATE	
	EPA		Plotida Target				:	EA-10		
Anlye	Meithed	MDL ⁶⁴	Chanup Concentration**	Units	(9/1/17)	P61713	\$	170796	96/8/11	1/10/97
Votatile Organic Aromatice										
Benzeise	07/18/109	0 6.1	_	l/an	•		-	,	`	7
Tolura	602/3030	-	. *	7/37	. 27	<u> </u>	, 0	7 7	7 7	, 7
Ethylbenian	602/8020	16.0	¥	7	210	360	\$	7 7	7 7	; 7
Xylana, Total	602/8010	0.9.3	< z	1/37	1,000	1 440	\$ 10 0	; ;	, v	; 7
Toul VOA	602/8010	0.0	2	ue/L	1.244	1.830	711	, V	7	; ,
Mechyl Ten-Bulyl Ether	209	1-25	: .	nr/l.	57.5		Ţ	7	7 7	; ;
1.3-Dieldomhensene	601/1010		××		7.	V	; ¬	7 7	7 7	7 7
1,4-Dichlerotenzencie	007/8/109	_	× Z	7/20	200	7.	7 7	; ;	, -	; 7
1,2-Dichlorobeazzae ⁴	602/8020	-	* X	1	3	\$	V	; -	, ,	7 7
Chlorobeniene	602/1010	1-1.3	YY	15	⊽	₹	v	~	⊽	⊽
Polynydrar Aromatic Hydracarbons		•								
Naphthaltrae	610/8100/8270	1-10	KX	₩5/L	62	96	190	1>	~	7
1-Methylauphiliatene	610/8100/8230	01-1	~Z	ug/L	!	i	3	~	_	7
2-Methyliaphiliatene	610/8100/8270	-10	42	1/3rd	11	i	24	7	~	J
Total Naphthalene	610/8100/8270	01-1	201	J/3H	95	g	111	~	v	~
Aceraphthytene	610/8100/8210	1.10	NOF	1/3m	01 >	2.5	4	v	~	⊽
Accruphitane	610/8100/8270	1.10	MDF	7/34	01×	~	011	7	7	~
Fluorene	610/8100/8210	01-1	MDL	7/84	o1 >	4.6	911	v	7	v
Thenothere	610/8100/8210	01-1	MDL	אמי	61 ×	2.1	53	~	-	~
2.4-Dimethylphenol	810W8270	2	<z< td=""><td>JOH T</td><td>2</td><td>:</td><td>:</td><td>:</td><td>:</td><td>:</td></z<>	JOH T	2	:	:	:	:	:
District, A) ambracenc	610/1100/1270	9:-	אומר	אפיר	91>	~	21.	⊽	~	⊽
Antiracese	610/8100/8370	1.10	NDF	7,84	DI >	ī	o1 >	7	<u>~</u>	⊽
Metadilotobuladiene	0153/0011	0	42	1/B/L	2 >	i	:	:	7	<u>-</u>
Yolulik Oreanic Halecerbous"										
Tetrachlarocthene	6010	-	4 2	7/31	•	:	:	:	!	:
1,4-Dikinlambenzene	9010	_	¥Z	1/3/	-	:	:	:	:	:
1,2-Dichlarobenzene	90108	_	۲	7/5/	-	:	!	!	į	:
l, J-Dichlatebenizat	0109	-	4	7/8d	~	:	:	1	1	:
1,2-Dhramoctbane	200	0.03	0.07	7/51	< 0 02	:	ı	i	÷	:
1,2-Dichlaracthaue	0101/109	1-5		1/4.1	2	:	*	į	:	:
Total Petraleum Hydrocarban	418.1	0.6-1	¥	mp/L	0.82	i	4	:	:	i
Ckad	139.1	2	\$1	Van	0	i	;	ļ	:	
		:	:			:	•	!	:	i

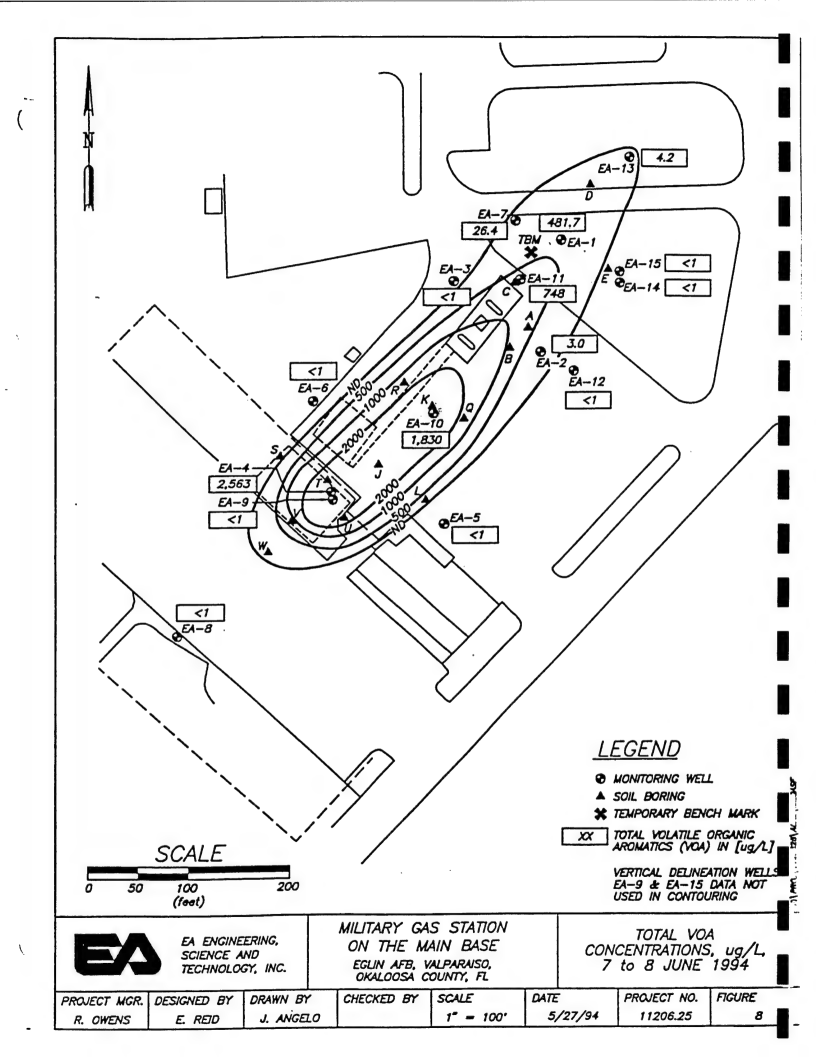
TABLE 4 (Cont.,

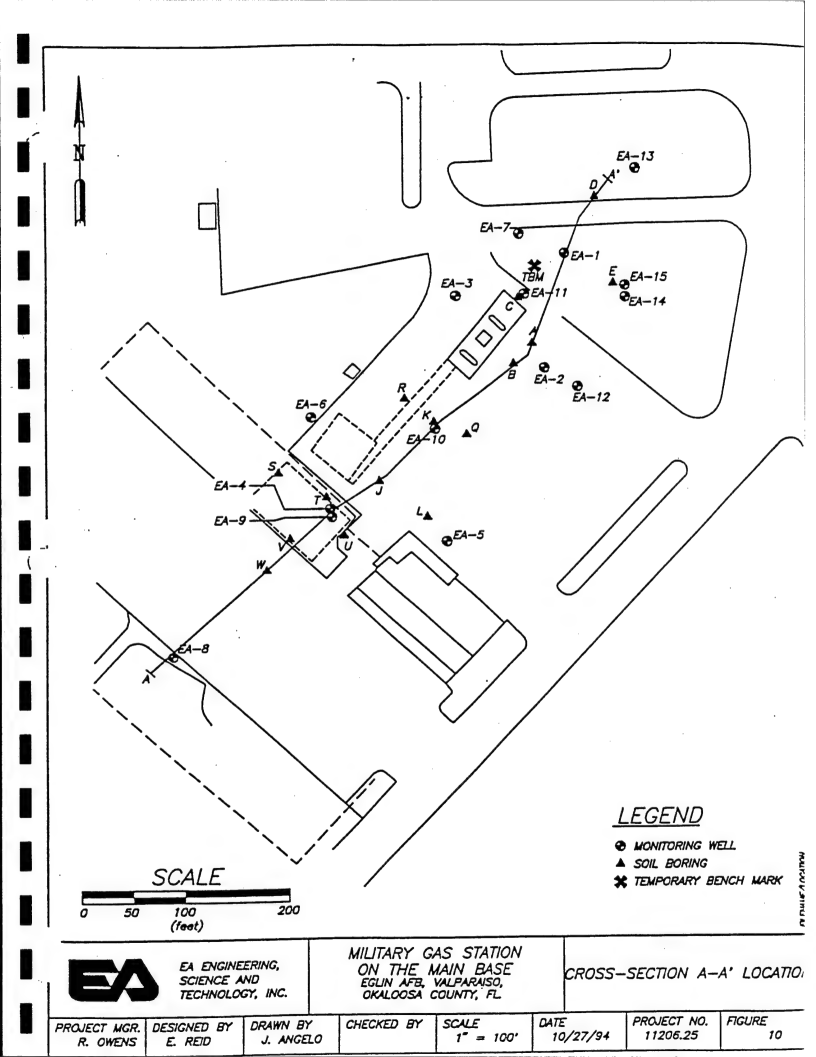
						WELL ID	WELL IDENTIFICATION AND SAMFLE DATE	AND SAMFLE	DATE	
	EPA		Florids Targer				EA-II	_		
Auslyte	Method	MDL*	Cleanup Concentration**	Units	1/1/43	6/7/94	; section)	1/20/96	94/3/11	2/10/97
Volsille Orennis Aromaiks"										
Benzene	0701/109	0 6.1	_	ייניור	₹	7	V	>	•	×
Toluene	602/1030	_	XX	1/34	Ţ	1	⊽	~	₹	7
Eulylbenrent	602/M120	1.6.0	NA NA	ושו	ı	8	1.3	4	5.3	7.8
Xylenes, Total	601/8010	11.9-2	٧٧	ייביר	62	487	⊽	7.0	5.4	3.0
Tetal VOA	601/8010	0.6-1	2	, F. M.	96	148	2.3	7.	10.7	K.S
Mechyl Ten-Buryl Ether	209	1-25	3	7/34	01>	:	~	•	J	⊽
1.3-Dichlesebenzene	602,1030		¥	PEN.	7	7	-	~	۲>	V
f.4. Dichturabenzenett	602/81120	-	× ×	12,	4	~	⊽	V	~	⊽
1,2.Dietilorobenzene	601/100	_	ž	7,24	53	~	~	⊽	ī	~
Chlorobeatene	601/1030		××	7/34	ī	·	7		1	~
Polynucical Arabatic Mediocarhomis										
Naphilistene	610/8100/8270	9:-	¥	7/27	43	3	33	1.1	5	**
1.Mahrlashilakac	610/8100/8370	9:	¥ 2		: !	: :	: 2	: =	7 7	2 0
2. Metty Insplit stone	610/8100/8270	9	× ×		-	: ;	: 2	: 3	7 7	
Total Naphthalene.	610/810Q/8170	1.0	98	7	: 5	3	2	24.1	= -	
Accesphihytene	610/8100/8270	==	NDF	7/27	01 >	3.8	- 4	2.1	; ·	
Aceasphilicine	610/8100/8170	1.10	MDL	7/8/	2 >	6.7	6.0	6.7	3 V	V
Fluorenc	610/810U/8270	<u> </u>	MDL	1/3rd	01×	7	71	-	د ا د	7
Plicasahiese	610/11/00/13/0	0:-	MDL	מניור	O1 >	~	1.5	-	<u>-</u>	~
2.4-Dimethylphenel	B100/8170	2	¥	re/	oi >	:		:	:	:
Dibent(1,6) inthrocme	610/8100/870	0:	MDL	7,31	01 >	~	1>	7	, † v	-
Anthracene	0128/0018/019	0:-	NOL	1/2/r	=>	~	7.6	⊽	3 ~	V
Heasthlorobutalisas	0/28/0018	9	¥	7,31	01 >	:	:	i	\$ - V	~
Yelallic Ottunk Helecarimmin										
Tetrachlorachene	202	_	٧2	1/5/1	-	:	:	:	:	:
1,4-Dichlorabenzene	202	-	Y X	I'd'I	~	:	:	:	:	:
1.2-Dichlorabenzene	1108	-	42	112.11.	⊽	:	:	:	:	:
I,3.Dichlorobenzene	9010	-	KY	7.21	⊽	:	i	!	:	:
1,2-Dibremocthane	\$	0.03	0 03	7/8/1	< 0.03	ı	į	:	1	:
1,2-Dkhlerotibene	0101/109	7.	n	HI/L	~	:	ŧ	:	÷	ı
Total Petraleum Elydrocarbon	418.1	0.6.1	¥X	7,210	> 0.6	•	ł	:	÷	:
Lead	239.2	2	. 21	784	v	:	i	i	:	:

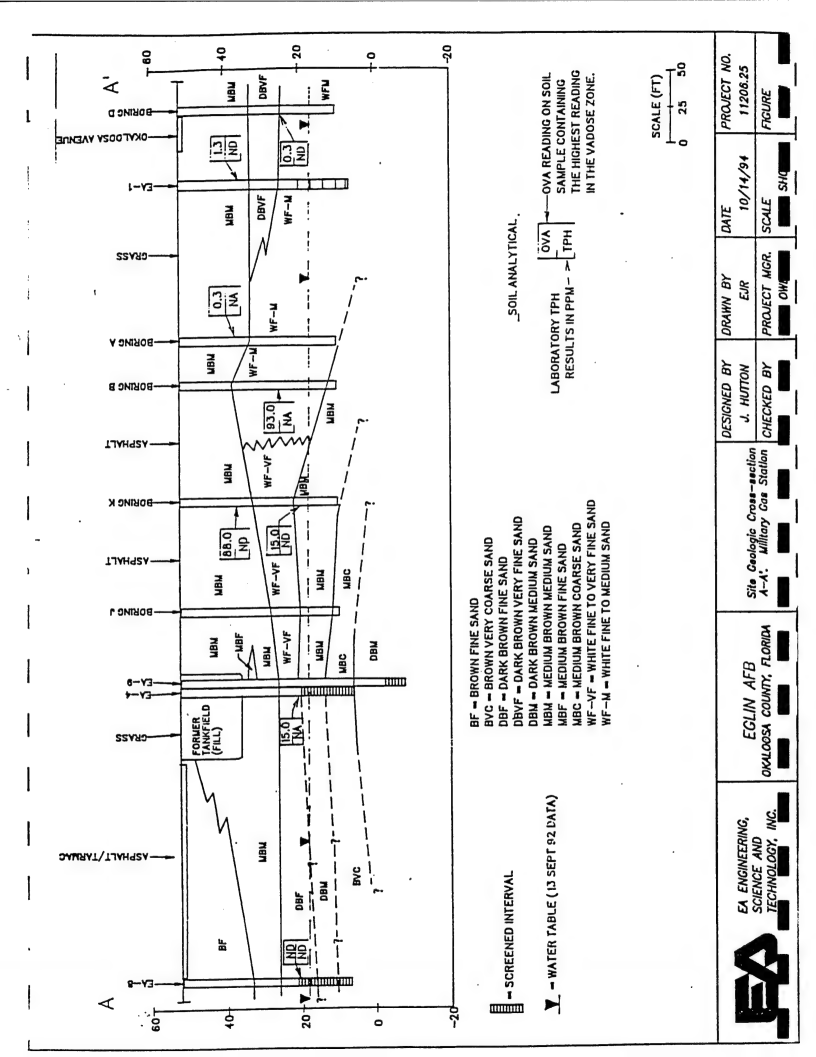
TABLE 4 (cont.)

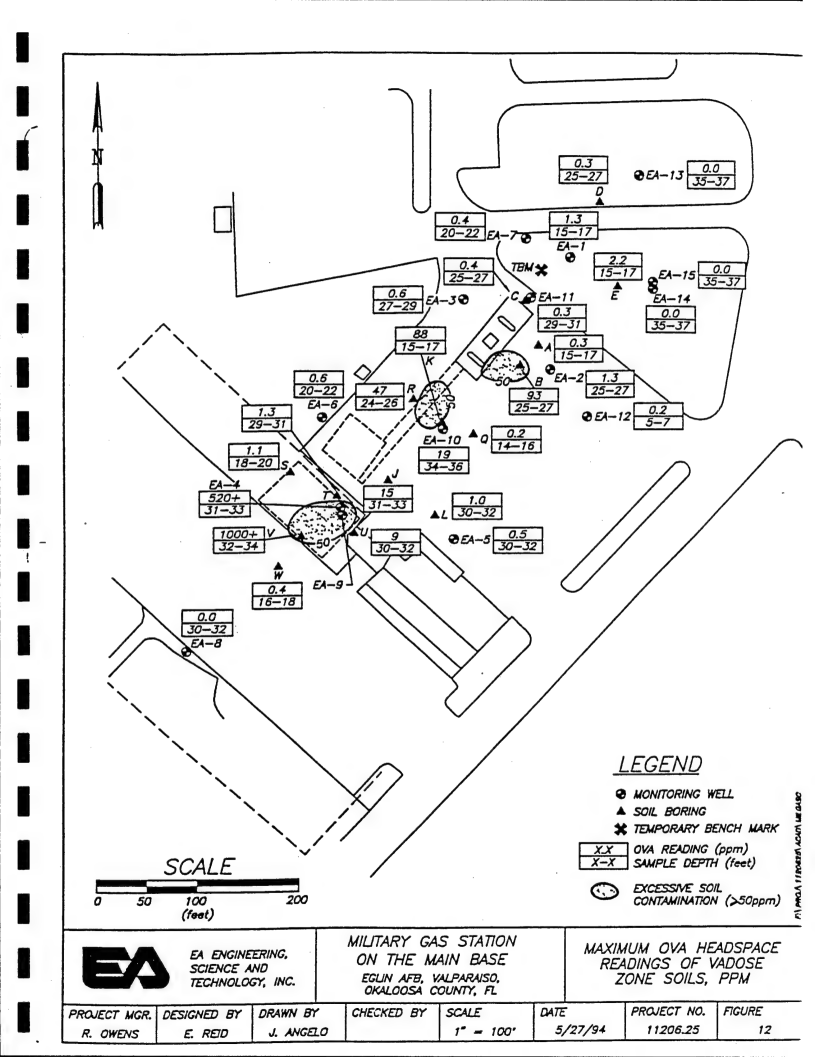
	· ·						WELL ID	WELL IDENTIFICATION AND SAMPLE DATE	AS GNA NC	WPLE DATE		
	EPA		Florida Targel		Ð	EA-12	Ē	EA-13	n,	EA-14		EA-15
Analyte	Number	MDC*	Concentation**	Units	103/94	67754	3/23/94	W17.00	3/23/94	50/05	3723/94	10/09
Velaille Organic Arematica"												
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Toluche	601/4010	-	ž	7,00	~	V	7	7	7 7	7	7	7
Ethylberaene	602/4020	_	¥	7	; 7	, ,	; ;	7:	7	7	<u>,</u>	⊽ '
Xyknez, Total	602/4020	_	¥		7	; ;	: 7	• •	7 7	7	₹ ;	₹ :
Total VOA	001/109	~	20		7 7	7	; ;	; ;	7 7	7	Ţ :	Ÿ :
Makyl Tent-Butyl Educr	603	_	9		; 7	;	; ;	7.	J ;	<u>-</u>	⊽ :	₹
1,3-Dichlerobenzeach	601/1010	_	2		7 7	: 7	7 7	; ;	,	: :	▼	i
1.4-Dichlotobeattach	601/1020	-	**	-	7 7	, ,	7 7	7	7	₹ :	<u>.</u>	₹ '
1,3-Dichlorabenzeoe**	602/8020	-	×		7 7	77	77	7	7	7	₹ :	⊽ :
Chlaratene	010\$/109	-	ž	ווי דיין	, ⊽	7 🗸	7 7	7 V	. .	⊽ ₹	⊽ ;	₹ ₹
Palanches Americal Madenastan								•	;	,	;	7
Naphhalene	45520012013	•	:									
1. Methyland Mariens	D/78/ODIBATIO	_	ž	77.	ī	-	~	~	7	•	•	~
2. McG with a labelle on	OTRANIBATIO	-	ž	727	!	1	* * *	!	1	1	:	: :
Total Nephibaleses	O/78/mile	:	ž	バブ	•	i	;	!	!	i	i	i
According	0/17/0018/019	_	8	161	<u>-</u>	⊽	1.3	~	•	7	1 >	>
Acranhibera	OUTRACIEMIE	_	MDL	ブ	⊽	ī	<u>~</u>	<u>~</u>	V	~	₹	V
Elizabeth Statement	O/TEMPIRALIO	-	HDF	121	₹	⊽	v	~		7	; 7	
בותוכנוני בייון	0/28/00/8/019	-	NDL	75.	~	7	<u>~</u>	~	7	7	7	7
ricesonaricos	610/11/0/11/0	-	NOL	PE/L	~	V	V	7	7	; ;	7 7	; ;
2.4.Dimethylphenol	6100/8270	:	ž	#1/L	į	i	:	; 1	; i	;	;	,
Dibeau(a,h)amhracene	610/8100/8170	-	MOL	ue/L	~	•	ī	ī	iī	: 7	: -	: ;
AMINITATION	610/8100/8170	-	MDL	7	~	V	· •	⊽	, ,	7 7	٠ ټ	7 7
Yalatlic Aceanis Ilabora bonz"	٠.										;	;
Tauchlerordune	601/100	!	¥	1700	;	i						
1,4-Dichlorobenrene	0101/109	!	¥		: !		!	:	:	:	i	:
1,2.Dichlorobenzene	0108/109	!	ž	100	! :		!!	i	:	;	:	:
1.2.01branderber							!	i	:	:	!	:
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1,2.Dichloroethane	0100	!	•	101	:	i	!	:	i	:	!	:
Total Petroleom Hydrocuebon	4IR.I	i	XX	mf/L	i	:	j	:	:	•	*	:
Lod	239.7		•									
	239.2	:	**	11-11								











APPENDIX C

FIELD DATA FORMS

Denver, Colorado

GS - Ground Surface

C - CORE

TOC - Top of Casing

G - GRAB

NS - Not Sampled

SAA - Same As Above U - Undetected

Water level drilled

Risk-Based Approach to Remediation

PARSONS ENGINEERING SCIENCE, INC.

Denver, Colorado

NOTES

30-

SAMPLE TYPE

bgs - Below Ground Surface

D - DRIVE

TD 29' has

GS - Ground Surface

C - CORE

TOC - Top of Casing

G - GRAB

NS - Not Sampled

U - Undetected

SAA - Same As Above

▼ Water level drilled

Risk-Based Approach to Remediation

GEOLOGIC BORING LOG

PARSONS ENGINEERING SCIENCE, INC.

Denver, Colorado

	Ę	Φ Bailer	made of:	dispos	able		
			type:describe:		ゼ		
	Sa	ample obta	ined is [X	G GRAB;	[] COM	POSITE SAMP	LE
ON-SI	TE MEA	SUREME	ENTS:			J	
Time							Measured wi
Temp (°	C)	25.9	25.0	26.1	26.0	26.0/	75 Z 55
pН		8.25	6.26	6.20	6.20	6.19	0111n 2501
Cond (µ		1130	1/33	1-33	1-33	1.32	Hach
DO (mg/		1.20	2.02	155	1.19	1.18	75255
Redox (n		-147.5	-171,1	-175.7	-179.4	-181.0	Oria 2501
gallons p	urged		5	6	7	8	
ON-SIT	E SAM	PLE TREA	ATMENT:	p · · · · ·			
	E SAM	- 7006	ATMENT:	od ## <i>Drss</i> od_			sound poly
ON-SIT	E SAM	PLE TREA	ATMENT: Meth Meth	od ## <i>Drss</i> od_		Containers:_	
ON-SIT	E SAM	PLE TREA	ATMENT: Meth Meth added: Meth	od F Orse od od	solur f less	Containers: Containers: Containers:	Sound poly 1-Liter and-15
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ON-SIT	E SAM	PLE TREA	ATMENT: Meth Meth added: Meth Meth Meth	od APOSS od_ od_ od_ od_TE/H od_STEX od_APOSS	ight for	Containers: Containers: Containers: Containers: Containers: Containers:	Sound poly 1-Liter and 1-15 40ml vial
ON-SIT	E SAM	PLE TREA	ATMENT: Meth Meth added: Meth Meth Meth	od Form	ight less	Containers: Containers: Containers: Containers: Containers: Containers: Containers:	Sound poly 1-Liter ambors 40ml vial 40ml vial
ON-SIT	E SAMI Fil	PLE TREA	ATMENT: Meth Meth Meth added: Meth Meth Meth Meth	od APOSS od_ od_ od_ od_TE/H od_STEX od_APOSS	ight less	Containers: Containers: Containers: Containers: Containers: Containers: Containers:	Sound poly 1-Liter and 1-15 40ml vial
ON-SIT	E SAMI Fil	PLE TREA tration: Servatives ANDLING	ATMENT: Meth Meth Meth Meth Meth Meth Meth Met	od Possod od Tepp od T	ight less	Containers: Containers: Containers: Containers: Containers: Containers: Containers:	Sound poly 1-Liter ambors 40ml vial 40ml vial
ON-SIT	E SAMI Fil	PLE TREA tration: Servatives ANDLING Contain Contain	ATMENT: Meth Meth Meth Meth Meth Meth Meth Met	od Ply od_ od_ od_ od_ od_ od_ od_ od_ od_ od_	lead ref	Containers: Containers: Containers: Containers: Containers: Containers: Containers:	Sound poly 1-Liter ambors 40ml vial 40ml vial
ON-SIT	E SAMI Fil	PLE TREA tration: Servatives ANDLING Contain Contain	ATMENT: Meth Meth Meth Meth Meth Meth Meth Met	od Possod od Tepp od T	lead ref	Containers: Containers: Containers: Containers: Containers: Containers: Containers:	Sound poly 1-Liter ambors 40ml vial 40ml vial
ON-SIT	E SAMI Fil Pre	PLE TREA tration: Servatives ANDLING Contain Contain	ATMENT: Meth Meth Meth Meth Meth Meth Meth Met	od Ply od_ od_ od_ od_ od_ od_ od_ od_ od_ od_	lrad	Containers: Containers: Containers: Containers: Containers: Containers:	1-Liter ambos 40ml vial 40ml vial 500 ml aclu

GROUNDWATER SAMPLING RECORD Eglin AFB F!

Sampling Location Homestead ARB Sampling Dates 3-16-97

GROUND W.	ATER SAMPLING RECORD - MONITORING WELL M: LA	- 4
		(Identification)
DATE AND SAMPLE CO	R SAMPLING: [X] Regular Sampling; [] Special Sampling; TIME OF SAMPLING: 3-26-99 at 14:45 a.m./fi.m. PLLECTED BY: Regular Sampling; [] Special Sampling; at 14:45 a.m./fi.m. Plant Clear Supply 2-75° F	
DATUM FOR	WATER DEPTH MEASUREMENT (Describe): 100 top	of Cosing
MONITORIN	IG WELL CONDITION:	
	[] UNLOCKED WELL NUMBER (3 - IS NOT) APPARENT STEEL CASING CONDITION IS: 0 -	
	INNER PVC CASING CONDITION IS: 04	
	WATER DEPTH MEASUREMENT DATUM (IS - IS NOT) APPAREN [] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR [] MONITORING WELL REQUIRED REPAIR (describe):	т
Check-off 1 []	EQUIPMENT CLEANED BEFORE USE WITH a corex di 150 Items Cleaned (List): a // in 5 trum core prob	di des
2[]	PRODUCT DEPTH	FT. BELOW DATUM
	WATER DEPTH 33.82 Measured with: Slage water ! puc! ; die flue	FT. BELOW DATUM
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe): Appearance: Clast Odor: 51.745 Other Comments:	·
4[]	WELL EVACUATION: Method:	

[] Bailer made of [] Pump, type: practifice [] Other, describe: Sample obtained is [X] GRAB; [] COMPOSITE SAMPLE ON-SITE MEASUREMENTS: Time Temp (°C) 25. F 250 25.1 25.0 455 55 pH 6.25 6.26 6.20 6.20 97.1 25.0 9			Monito	Groundwater	Sampling R					
Pump, type:	5[]	SAMPLI	E EXTRACTION	METHOD:	-/7 7	_ (Continued)				
Time Temp (°C) 25- K 250 25.1 25.0 Measured with Temp (°C) 25- K 250 25.1 25.0 VSC 55 pH 6.25 6.26 6.29 6.21 00.1.256/2 Cond (µS/cm) -129 -170 -190 -1/3 + trech DO (mg/L) 1.21 1.20 1.20 1.20 VSC 55 Redox (mV) -163.1 -110 -1/1.3 -171.4 grant 25.69 gallons purged 1 2 3 VSC 55 gallons purged 1 2 3 VSC 55 SAMPLE CONTAINERS (material, number, size): 9-wm class vials y-11.1-1 glass ON-SITE SAMPLE TREATMENT: [] Filtration: Method Containers: Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Containers			[] Bailer n [] Pump, t [] Other, o	nade of:	rc					
Time Temp (°C) 25- K 250 25.1 25.0 Measured with Temp (°C) 25- K 250 25.1 25.0 VSC 55 pH 6.25 6.26 6.29 6.21 00.1.256/2 Cond (µS/cm) -129 -170 -190 -1/3 + trech DO (mg/L) 1.21 1.20 1.20 1.20 VSC 55 Redox (mV) -163.1 -110 -1/1.3 -171.4 grant 25.69 gallons purged 1 2 3 VSC 55 gallons purged 1 2 3 VSC 55 SAMPLE CONTAINERS (material, number, size): 9-wm class vials y-11.1-1 glass ON-SITE SAMPLE TREATMENT: [] Filtration: Method Containers: Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Containers			Sample obtain	ined is TVI GDA	P. [] CO)	DOCUME CALL				
Time Temp (°C) 25. K 250 25.1 25.0	[]	ON-SITE	ON-SITE MEASUREMENTS:							
PH		Time					Managed with			
Cond (µS/cm)				25,0 25.1	25.0					
DO (mg/L) 1.21 1.20										
Redox (mV) -163.1 - 170 -171.3 -171.4 gass containers: SAMPLE CONTAINERS (material, number, size): 9-wm/ gass containers: [] Filtration: Method Containers: Method Containers: [] Preservatives added: Method Try Containers: // Containers: // Containers: // Method Containers: // Containers: // Containers: // Containers: // Method Containers: // Containers: // Containers: // Method Containers: // Containers: // Containers: // Containers: // Method Containers: // Conta										
gallons purged 1 2 3										
SAMPLE CONTAINERS (material, number, size): 9-wm class vials 4-11. 1-1 class ON-SITE SAMPLE TREATMENT: [] Filtration: Method Containers: Method Containers: Method Containers: Method Containers: Womloaks Nec Method Containers: Womloaks Nec Method Containers: Womloaks Nec Method Containers: Womloaks Nec Method Containers: Method Containers: Method Containers:					.3 -171.4		Orion 250A			
ON-SITE SAMPLE TREATMENT: [] Filtration: Method Containers: Method Containers: Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Method Containers: Contain		ganons pur	gcu							
Method Fer-Methone Containers: Vumluials Here Method Ir h Containers: /Li here glass Method Containers: Method Containers: Containers:		[]	Filtration:	Method		Containers:				
Method Containers: /Li Ley glass Method Containers: Containers: Method Containers: Containers		[]	Preservatives a	added:						
Container Sides Labeled Container Lids Taped Containers Placed in Ice Chest		·		Method_lr, A Method_	-Methode	Containers: //_: Containers:	moviels HEL For glass			
Container Lids Taped Containers Placed in Ice Chest]	CONTAIN	ER HANDLING	:						
/			[] Containe	er Lids Taped	Chest					
	1	OTHER CO	/							
		-								

GROUNDWATER SAMPLING RECORD

Eqlin AFB Milgas fation

Sampling Location Homestead ARB

Sampling Dates 7-31-98

GROUND W.	ATER SAMPLING RECORD - MONITORING WELL EA-4 CUP EA-4
REASON FO DATE AND 'SAMPLE CO WEATHER:	(Identification) R SAMPLING: [X] Regular Sampling; [] Special Sampling; I'IME OF SAMPLING: 3-31-98 at 0 1/5 a.m./p.m. LLECTED BY: 100 of Parsons ES WAIM 25/4 Same 7 R WATER DEPTH MEASUREMENT (Describe): 100 Top of resides
MONITORIN	WELL CONDITION: [DLOCKED: WELL NUMBER (IS - IS NOT) APPARENT STEEL CASING CONDITION IS: INNER PVC CASING CONDITION IS: WATER DEPTH MEASUREMENT DATUM (IS - IS NOT) APPARENT [] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR [] MONITORING WELL REQUIRED REPAIR (describe):
Check-off	EQUIPMENT CLEANED BEFORE USE WITH a canox di isa di Items Cleaned (List): all instrument probes
2[]	PRODUCT DEPTH
	WATER DEPTH 33 81' FT. BELOW DATUM Measured with: 5/ept wate level is helder
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe): Appearance:
4[]	WELL EVACUATION: Method: Volume Removed: Volume Remove

	EXTRACTIO		,	11		
	Bailer Pump	r made of:	refesal	512		
	[] Other	, type <u>rassa</u> , describe:_	-TILT FE		Pe-	
	[] 05.01	, accinoc				
	Sample ob	tained is [2	X] GRAB;	[] CON	POSITE SAME	PLE
ON SITTE	MEASUREM					
ON-SITE	MEADUREM	ENIS:				
Time						Measured
Temp (°C)	22-6	23.0	22.8	23.0	23.0	Measured
pН	6.75	6.63	6.27	6-34	6,33	
Cond (µS/c		1./16	.122	1119	,120	
DO (mg/L)	6.31	6.03	6.30	6.37	6.02	
Redox (mV)		67.3	73.6	64.6	64.7	
gallons purg	ged 0		12	3	4	
	SAMPLE TRE					
		ATMENT:	iod		Containers:	
ON-SITE S	SAMPLE TRE	ATMENT: Meth Meth	nod		Containers: _ Containers:	rials, 1-14it
ON-SITE S	SAMPLE TRE	ATMENT:	nod		Containers: _ Containers:	
ON-SITE S	SAMPLE TRE	ATMENT: Meth Meth Meth	nod		Containers: _ Containers:	
ON-SITE S	SAMPLE TRE	ATMENT: Meth Meth Meth Sadded:	nod_ nod_ nod_		Containers: Containers: Containers:	
ON-SITE S	SAMPLE TRE	ATMENT: Meth Meth Meth s added: Meth Meth	od_ od_ od_ od_ od_ od_ od_ od_ od_	~ 6	Containers: Containers: Containers:	youl vial
ON-SITE S	SAMPLE TRE	Meth Meth Meth s added: Meth Meth Meth	od_od_od_od_od_od_od_od_od_od_od_od_od_o		Containers: Containers: Containers: Containers: Containers:	youl vial
ON-SITE S	SAMPLE TRE	Meth Meth Meth s added: Meth Meth Meth	od_ od_ od_ od_ od_ od_ od_ od_ od_		Containers: Containers: Containers: Containers: Containers:	you I via!
ON-SITE S	SAMPLE TRE	Meth Meth Meth s added: Meth Meth Meth	od_od_od_od_od_od_od_od_od_od_od_od_od_o		Containers: Containers: Containers: Containers: Containers: Containers:	you I via!
ON-SITE S	Filtration: Preservative	Meth Meth Meth Sadded: Meth Meth Meth Meth	od_ od_ od_ od_ od_ od_ od_		Containers: Containers: Containers: Containers: Containers: Containers:	you I via!
ON-SITE S	Filtration: Preservative ER HANDLIN [N] Conta	Meth Meth Meth s added: Meth Meth Meth Meth Meth Meth	od_sod_sod_sod_sod_sod_sod_sod_sod_sod_s	~ <	Containers: Containers: Containers: Containers: Containers: Containers:	you I via!
ON-SITE S	Filtration: Preservative ER HANDLIN [N] Conta	Meth Meth Meth s added: Meth Meth Meth Meth Meth Meth	od_sod_od_od_od_od_od_od_od_od_od_od_od_od_o	~ <	Containers: Containers: Containers: Containers: Containers: Containers:	you I via!
ON-SITE S	Filtration: Preservative ER HANDLIN [] Conta [] Conta	Methods added: Methods added:	od_bod_od_od_od_od_od_od_od_od_od_od_od_od_o	est est	Containers: Containers: Containers: Containers: Containers: Containers:	you! via!

GROUNDWATER SAMPLING RECORD Elgin AFB F1.

Sampling Location Homestead ARB
Sampling Dates 3-27-99

GROUND W	VATER SAMPLING RECORD - MONITORING WELL M; / EA - 5
	(Identification)
DATE AND SAMPLE CO	OR SAMPLING: [X] Regular Sampling; [] Special Sampling; TIME OF SAMPLING: 3-27-98 at 0930 for L/p.m. OLLECTED BY: \(\frac{\partial 0}{\partial 0} \) of \(\partial 0
DATUM FO	R WATER DEPTH MEASUREMENT (Describe): Toc Top of casing
-	
MONITORI	NG WELL CONDITION:
	[/] LOCKED: [·] UNLOCKED
	WELL NUMBER (IS - IS NOT) APPARENT
	STEEL CASING CONDITION IS: 0 K
	WATER DEPTH MEASUREMENT DATUM (IS - IS NOT) APPARENT
	DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR
•	[] MONITORING WELL REQUIRED REPAIR (describe):
Check-off	
1[]	EQUIPMENT CLEANED BEFORE USE WITH Alconox, Di, Iso, Di. Items Cleaned (List): 2// justrument probs
	Items Cleaned (List): all justrument probs
2[]	PRODUCT DEPTH hold FT. BELOW DATUM
	Measured with: Made! 121 interface meter
	WATER DEPTH 33.81 FT. BELOW DATUM
	WATER DEPTH 33.81 FT. BELOW DATUM Measured with: 5/ape water level indicator
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe): Appearance: Clad
	Odor: pon C
	Other Comments:
4[]	WELL EVACUATION:
4[]	Method: pure?
	Volume Removed: ~ 3 a a /
	Observations: Water (slightly - very) cloudy e (car
	Water level (rose - fell no change) Water odors: Man
	Other comments:

Groundwater Sampling Record Monitoring Well No. En - 8 (Continued) 5[] SAMPLE EXTRACTION METHOD: M Bailer made of: dergoschle (goly) [] Pump, type:_ [] Other, describe: Sample obtained is [X] GRAB; [] COMPOSITE SAMPLE 6[] ON-SITE MEASUREMENTS: Time Measured with Temp (°C) 24.0 240 24.3 24.3 YSI 55 pН 6,16 6.27 6.27 6.27 Orion 250p Cond (µS/cm) 096 .102 100 .102 .102 Hach DO (mg/L) 7.26 6.18 6.15 6.08 6.11 451 55 Redox (mV) 98.6 94.5 93.2 79.1 90.1 Orion 250x gallons purged SAMPLE CONTAINERS (material, number, size): 9-40ml glass Links 1-12:ter glass 7[] 8[] ON-SITE SAMPLE TREATMENT: [] Filtration: Method Containers: Method Containers: Method Containers: [] Preservatives added: Method BTEX Containers: 40ml vial HCL Method Mother e Containers: 40 al vial Method Containers: Method Containers: 9[] CONTAINER HANDLING:

Container Sides Labeled

Containers Placed in Ice Chest

Container Lids Taped

K

OTHER COMMENTS:_

c:\admin\gwsample.doc

10[]

GROUNDWATER SAMPLING RECORD ELLIN AFB F1.

Sampling Location Homestead ARB
Sampling Dates 3-27-52

GROUND Y	WATER SAMPLING RECORD - MONITORING WELL M. 1 EA - 8	
	(Id	lentification)
REASON F	OR SAMPLING: [X] Regular Sampling; [/] Special Sampling;	
DATE AND	TIME OF SAMPLING: 3-27-97 at 1/3 0 a.m./p.m.	
SAMPLE C	COLLECTED BY: LOS of Parsons ES	
WEATHER	· Warm Gunge - 7/0F	
DATUM FO	DR WATER DEPTH MEASUREMENT (Describe): 700 700	Casing
MONITOR	ING WELL CONDITION:	
	[A] LOCKED: [] UNLOCKED	
	WELL NUMBER (IS) IS NOT) APPARENT	
	STEEL CASING CONDITION IS: 6K	
	INNER PVC CASING CONDITION IS: 62	
	WATER DEPTH MEASUREMENT DATUM (S- IS NOT) APPARENT	
	DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR	
	MONITORING WELL REQUIRED REPAIR (describe):	
	[] MONTOIGNO WEDE ICEQUIRED ICE AIR (GOOTIOS).	
Check-off		
1[]	FOURMENT OF FANED REFORE USE WITH A CAPARA L.	/.·
- L J	EQUIPMENT CLEANED BEFORE USE WITH a conexide, iso, of Items Cleaned (List): all instrument from	
	Homs Cleaned (BBI). (EV. / A) / [EA 2]	
2[]	PRODUCT DEPTH None	FT. BELOW DATUM
	Measured with: Motol 121 onto face motor	
	WATER DEPTH Measured with: Slope water level indicate	FT. BELOW DATUM
	Measured with: Stope water touch indicate	
3 []	WATER-CONDITION BEFORE WELL EVACUATION (Describe):	
	Appearance: C/rar	
	Odor: Mone	
	Other Comments:	
4 []	THELL EXACTIATION.	
4[]	WELL EVACUATION:	
	Method: pump Volume Removed: ~ V 5a/	
	Observations: Water (slightly - very) cloudy c'res	
	Water level (rose - fell no change)	
	Water odors: Non C	
	Other comments:	

	Cample shee	describe:_				
ON-SITE M	EASUREME		G GRAB;	[] COM	POSITE SAMPI	LE
Time						Measured v
Temp (°C)	24.8	24.4	23.9	24.6		45t 55
pН	762	7.58	7.54	7.54		Orien 25
Cond (µS/cm		./35	1./35	./35		Hach
DO (mg/L)	7.50	6.56	7.90	7.5/		45I 55
Redox (mV) gallons purge	61.1	68.5	34-7	34.5		0, 50
ON-SITE SA	ONTAINERS 2-500	ATMENT:			•	15, 5-1-Liter
ON-SITE SA	ONTAINERS 2-500	ATMENT: Meth Meth			•	Soon poly
ON-SITE SA	ONTAINERS 2-500	ATMENT: Meth Meth	nod <u>disell</u>		Containers:	Soon poly
ON-SITE SA	ONTAINERS 2 - 500	ATMENT: Meth Meth sadded: Meth Meth Meth	and disself and and and and and and and and and and	est	Containers: Containers: Containers: Containers: Containers: Containers: Containers:	SODAL paly Youl vial Loal vial L-Liter ambor
ON-SITE SA	ONTAINERS 2-500 AMPLE TREA Filtration: Preservatives	ATMENT: Meth Meth Meth Meth Meth Meth Meth	and sell	est	Containers: Containers: Containers: Containers: Containers: Containers: Containers:	SODMI paly

GROUNDWATER SAMPLING RECORD Eglin AFB FL. Sampling Location Homestead ARB

Sampling Dates 3-27-ay

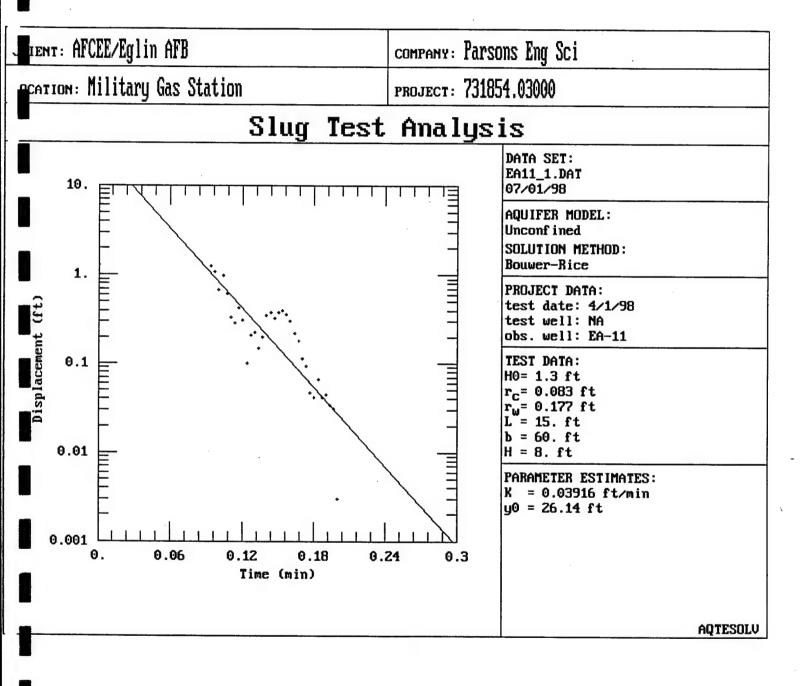
GROUND W	VATER SAMPLING RECORD - MONITORING WELL Mil EA-	10
DATE AND	OR SAMPLING: [X] Regular Sampling; [] Special Sampling; TIME OF SAMPLING: 3-27-9/ at 0730 (a.m./p.m. OLLECTED BY: ROLL OF Parsons ES Usen, Sunny ~ 620 F	f casia 5
MONITORI	WELL CONDITION: [A] LOCKED: WELL NUMBER (IS) IS NOT) APPARENT STEEL CASING CONDITION IS: INNER PVC CASING CONDITION IS: WATER DEPTH MEASUREMENT DATUM (IS) IS NOT) APPARENT [] DEFICIENCIES CORRECTED BY SAMPLE COLLECTOR [] MONITORING WELL REQUIRED REPAIR (describe):	•••
Check-off 1 []	EQUIPMENT CLEANED BEFORE USE WITH a leaner de ise, de Items Cleaned (List): all instrument quality	
2[]	PRODUCT DEPTHnone	FT. BELOW DATUM
3[]	WATER-CONDITION BEFORE WELL EVACUATION (Describe): Appearance: Class Odor: Mana	
4[]	WELL EVACUATION: Method:	

APPENDIX D SLUG TEST DATA ANALYSIS

CLIENT: AFCEE/Eglin AFB company: Parsons Eng Sci LOCATION: Military Gas Station РВОЈЕСТ: 731854.03000 Slug Test Analysis DATA SET: EA10_1.DAT 07/01/98 AQUIFER MODEL: Unconfined SOLUTION METHOD: Bouwer-Rice PROJECT DATA: Displacement (ft) 1. test date: 3/30/98 test well: NA obs. well: EA-10 TEST DATA: H0 = 2.5 ft $r_{c} = 0.083 \text{ ft}$ $r_{W} = 0.177 \text{ ft}$ L = 15. ft 0.1 b = 60. ftH = 9. ftPARAMETER ESTIMATES: K = 0.01038 ft/miny0 = 54.88 ft0.01 0.4 0.8 1.6 Time (min)

AQTESOLV

CLIENT: AFCEE/Eglin AFB company: Parsons Eng Sci LOCATION: Military Gas Station рвојест: 731854.03000 Slug Test Analysis DATA SET: EA10_2.DAT 07/01/98 AQUIFER MODEL: Unconfined SOLUTION METHOD: Bouwer-Rice PROJECT DATA: Displacement (ft) 0.1 test date: 3/30/98 test well: NA obs. well: EA-10 TEST DATA: H0= 0.8 ft r_c= 0.083 ft $r_{\mu} = 0.177 \text{ ft}$ 0.01 L = 15. ft b = 60. ft H = 9. ftPARAMETER ESTIMATES: $K = 0.02047 \, \text{ft/min}$ y0 = 0.7694 ft0.001 0.06 0.12 0.24 0.3 Time (min)



APPENDIX E

CALCULATIONS

Pore Volume calculation

Source Area length 50 ft.

Groundwater Velocity 124 ft/yr

50 ft = 0.4 for 1 pore Volume to migrate
124 ft/yr through the source area.

... 2,5 pore volumes /

.. 2,5 pore volumes/yr

Find t.

C= 400 mg/L (Xylenes @ EA-4, 1998)

C= 20 mg/L (FDEP TCL for Xylenes)
-k=-0.001/day

20 ng/L = 400 ng/L × e (0.001t/day)

0.05 = -0.001t/day

 $ln(0.05) = \frac{-0.001t}{day}$ -3 = -0.001 t

> 2996 day = t t = 8.2 yr

> > Client AFCEE Mil Gas Ealin AFB Proj. No. 73/854, 03000, 45.01, 3000 By Cindy Nagel Date June 25 1998 Chkd. Was 3000 Date 6 25/98

Decay Calc $C = C_0 e^{(-kt)}$

	Xylenes		Eth	nylbenze	ene	N	aphthalei	ne
C =	20	μg/L	C =	30	μ g/L	C =	20	$\mu g/L$
$C_0 =$	400	$\mu g/L$	$C_0 =$	76	$\mu g/L$	$C_0 =$	40	$\mu \text{g/L}$
k =	-0.0009	day-1	k =	-0.001	day-1	k =	-0.0007	day-1
t =	????	days	t =	????	days	t =	????	days
				000	•		222	
t =	3329	days	t =	930	days	t =	990	days
t =	9.1	years	t =	2.5	years	t =	2.7	years

Client Esta AFB	Job No. <u>731854, 03000</u>	Sheet_' of
Subject Xylene Volumes	By Steve Hicks	Date 3/8/99
Subject Afterne Holowica	Checked	Rev
200+ Area = 0,8" × 0,35" = 64' × 28' = 1792ft=	1:v = 20tt	
20+ Area = 2.15"x0.8"=173'x64'= 11,008ft2	···. · · · · · · · · · · · · · · · · ·	
5+ Area = 3,5" x 2"= 280' x 160' = 44,800f+		
200+ vol = 17929+2×59+ = 89609+3		
20+ Vol = 11,008 ft > x8ft = 88,064ft 3 (Includes	200+ Volume)	
5+ Nal = 44,800f+2×10f+ = 448,000f+3 (Includes	200+ and 20+ Volume)	
Actual Volume of soil and water		
5+ Vol = 448,000ft3-88,064ft3 = 359,936f	f_3	
90+No1 = 28'00Att3-8200tt3 = 20'10Att3		
200-101 = 8960++3		
Actual Volume of Worter fine sand, \$ = 30%		
5+Vol = 359,936ft3x 30% = 107,980,8ft3.	= 3,057676 L = 5+ Vol	
20+101 = 79,104 ft3 x 30% = 23,731,2ft3		
200+101 = 8,960 ft3 x 30% = 3,688 ft3 =	76,116L = 200+Vol	
1/4)		
Volume of Contaminant 5+ Vol = 3,057,676 L x 10 Mg/L = 30,5	7 760,0	
20+Vol = 671,993 L × 100 M3/L = 67,1	991300 Ma	
200+vol = 76,116L x 300 ms/L = 22,8	34,800 ug	
	10.860, mg = 120, 611 mg = 120.0	6 3
120.65 × 0.95 = 134 mL Xylene = 9.1	16 Hbsp	
		1

PESI-COR-8 (12/94)